



Kongeriget Danmark

Patent application No.:

1156/96

Date of filing:

18 Oct 1996

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TAASTRUP 09 Aug 1997

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NOVEL PROBES FOR THE DETECTION OF MYCOBACTERIA OF THE MYCOBACTERIUM TUBERCULOSIS COMPLEX

The present invention relates to the design, construction and use of novel probes for detecting mycobacteria of the Mycobacterium tuberculosis Complex (MTC) which probes are capable of detecting the organisms in test samples, e.g. expectorates, sputum, aspirates, urine, blood and tissue sections, food, soil and water.

BACKGROUND OF THE INVENTION

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Tuberculosis caused by mycobacterial infection is presently the predominant infectious cause of morbidity and mortality world-wide, and is estimated to kill about three million people annually. WHO estimates that the annual number of new cases of tuberculosis will increase from 7.5 million in 1990 to 10.2 million in 2000, an escalation that will result in approximately 90 million new cases during this decade. It is furthermore estimated that 30 million people will die from tuberculosis during the 1990s, which equals one quarter of preventable deaths among adults.

The prevalence of tuberculosis has been very high in the poorer parts of the world such as

Asia, Africa and South-America, but in recent years an increase has also been observed in industrialised countries. This appears to be due to an interaction of various factors including i.a. patterns of migration, poorly organised tuberculosis programmes and nutrition problems. Furthermore, a serious threat will arise from the emergence of new strains that are multi-drug resistant.

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Considering the perspective and impact the disease has, the development of rapid, specific and preferably easy-performed and economic feasible diagnostic detection tests are of utmost importance and would be a very valuable tool in the fight against the spread of tuberculosis.

Generally, mycobacterial infections are divided into infections caused by two groups of bacteria, namely mycobacteria of the Mycobacterium tuberculosis Complex (MTC) and mycobacteria of the Mycobacterium avium-intracellulare Complex (MAC). The mycobacteria of the Mycobacterium tuberculosis Complex include M. tuberculosis, M. bovis and M. africanum, whereas the mycobacteria of the Mycobacterium avium-intracellulare Complex comprise M.

avium and M. intracellulare.

Presently, the detection of mycobacteria by microscopy gives the more accurate diagnosis. The sample (e.g. an expectorate) is stained for the presence of acid-fast bacillus using Ziehl-

Neelsen staining and may subsequently be cultured in order to confirm the result obtained by staining. Such techniques are one of the cornerstones of all anti-tuberculosis programmes. However, the Ziehl-Neelsen staining lacks sensitivity since the detection limit is 10⁴ organisms/ml or greater. On the contrary, cultivation is sensitive, and it may be possible to detect 10-100 organisms per sample, but the result is not available before up to 8 weeks of cultivation. Likewise, information of drug susceptibility is not available until after 2-3 weeks of further testing.

Automated detection is rapidly becoming available for large scale testing for the presence of mycobacteria. Such systems include ESP Myco Culture System (Difco), MB/BacT (Organon Teknika) and MGIT (Becton Dickinson). These test methods are based on colorimetric or fluorometric detection of carbon dioxide or oxygen produced by mycobacterial metabolism.

Neither staining nor cultivation methods allows distinction between the mycobacteria of the MTC and the MAC.

Some of the attempts to replace the methods based on cultivation rely on target amplification or target hybridisation using specific probes.

One of such newly developed target amplification method is based on PCR. The principle of this reaction is, through amplification of specific nucleic acid sequences of the mycobacteria, to increase the copy number of the specific sequence to a level where it may be detectable in an early stage of the infection. In principle, the PCR reaction offers the possibility of detecting as few as one target sequence. In most cases, the DNA is extracted prior to carrying out the PCR reaction. However, it has become clear that the method used to extract DNA from specimens has a great influence on the sensitivity and specificity of PCR products.

Furthermore, false negative results in specimens may be obtained due to the presence of inhibitors of the PCR reaction such as haemoglobin and proteins.

Another problem arises from cross-contamination of negative specimens with a bacteria not present in the sample. This may cause problems in conventional bacteriological procedures and may lead to a positive PCR result. Contamination of reagents and specimens with amplified PCR products is yet another well-recognised problem when using a PCR-based

diagnosis.

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Nucleic acid probes for detecting rRNA of mycobacteria have been described in for example US 5 547 842, EP-A 0 572 120 and US 5 422 242.

SUMMARY OF THE INVENTION

The present invention discloses and claims novel peptide nucleic acid probes for the detection of mycobacteria of the Mycobacterium tuberculosis Complex. The probes detect sequences in 23S rRNA and genomic sequences corresponding to said rRNA. rRNA is present in a high number of copies in each cell, and hence a well suited target for a sensitive test. Furthermore, probes that are complementary to said rRNA are especially suitable for hybridisation as it is known that species variable regions exist within these highly conserved sequences thereby enabling the design of probes for detecting mycobacteria of the Mycobacterium tuberculosis Complex.

The novel probes may be used in an assay for the detection of mycobacteria of the MTC group. The mycobacteria of the MTC group are responsible for significant morbidity and mortality in humans. M. tuberculosis is the most common mycobacteria of the MTC group isolated from humans. M. bovis may be transmitted from infected animals to humans. M. africanum causes pulmonary tuberculosis in tropical Africa.

Tuberculosis is highly contagious, and a rapid diagnosis of the disease is therefore very important. For most clinical laboratories, assignment of an isolate to the group of MTC bacteria is sufficient.

Thus, in a first aspect, the invention features a hybridisation assay probe able to detect mycobacteria of the MTC group. Specifically, the probe is a peptide nucleic acid as defined in claim 1. Such probes do not to any significant degree cross react with nucleic acid from other organisms in the test sample under appropriate stringency conditions.

In another aspect, the present invention relates to a method according to claim 7 for detecting the presence of organisms belonging to the MTC group.

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In yet another aspect, the present invention relates to a kit comprising at least one peptide nucleic acid probe as defined in anyone of claims 1 to 6.

BRIEF DESCRIPTION OF THE FIGURES

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Figure 1 shows positions 281 to 400 and 601 to 680 of an alignment of 23S rRNA of M. tuberculosis (positions 1251 to 1368 and 1568 to 1647 of GenBank entry GB:MTCY130, accession number Z73992), M. avium (positions 579 to 697 and 896 to 960 of GenBank entry

GB:MA23SRNA, accession number X74494), M. phlei (positions 665 to 783 and 978 to 1047 of GenBank entry GB:MP23SRNA, accession number X74493), M. leprae (positions 591 to 709 and 909 to 973 of GenBank entry GB:ML5S23S, accession number X56657), M. paratuberculosis (positions 579 to 697 and 896 to 960 of GenBank entry GB:MPARRNA, accession number X74495), M. gastri (positions 366 to 484 and 684 to 748 of GenBank entry GB:MG23SRRNA, accession number Z17211) and M. kansasii (positions 309 to 427 and 627 to 691 of GenBank entry GB:MK23SRRNA, accession number Z17212). Preferred nucleobase sequences of probes may be chosen within positions 326 to 359 or 635 to 658 of the alignment corresponding to positions 1296 to 1327 and 1602 to 1625 of M. tuberculosis sequence (indicated by heavy frames). Mismatches between the sequences of M. avium, M. phlei, M. leprae, M. paratuberculosis, M. gastri and M. kansasii and that of M. tuberculosis in the alignment are indicated by light frames.

Figure 2 shows positions 721 to 840 of an alignment of 23S rRNA of M. tuberculosis (positions 1688 to 1807 of GenBank entry GB:MTCY130, accession number Z73992), M. avium (positions 1001 to 1109 of GenBank entry GB:MA23SRNA, accession number X74494), M. phlei (positions 1088 to 1203 of GenBank entry GB:MP23SRNA, accession number X74493), M. leprae (positions 1014 to 1125 of GenBank entry GB:ML5S23S, accession number X56657), M. paratuberculosis (positions 1001 to 1109 of GenBank entry GB:MPARRNA, accession number X74495), M. gastri (positions 789 to 900 of GenBank entry GB:MG23SRRNA, accession number Z17211) and M. kansasii (positions 732 to 843 of GenBank entry GB:MK23SRRNA, accession number Z17212). Preferred nucleobase sequences of probes may be chosen within positions 761 to 787 of the alignment corresponding to positions 1728 to 1754 of M. tuberculosis sequence (indicated by a heavy frame). Mismatches between the sequences of M. avium, M. phlei, M. leprae, M. paratuberculosis, M. gastri and M. kansasii and that of M. tuberculosis in the alignment are indicated by light frames.

Figure 3 shows positions 1281 to 1360 and 1601 to 1680 of an alignment of 23S rRNA of M. tuberculosis (positions 2246 to 2323 and 2563 to 2636 of GenBank entry GB:MTCY130, accession number Z73992), M. avium (positions 1549 to 1626 and 1865 to 1938 of GenBank entry GB:MA23SRNA, accession number X74494), M. phlei (positions 1643 to 1720 and 1960 to 2027 of GenBank entry GB:MP23SRNA, accession number X74493), M. leprae (positions 1565 to 1644 and 1884 to 1959 of GenBank entry GB:ML5S23S, accession number X56657), M. paratuberculosis (positions 1549 to 1626 and 1865 to 1938 of GenBank entry GB:MPARRNA, accession number X74495), M. gastri (positions 1339 to 1406 and 1646 to 1719 of GenBank GB:MG23SRRNA, accession number Z17211) and M. kansasii (positions 1282 to 1349 and 1589 to 1663 of GenBank entry GB:MK23SRRNA, accession number

Z17212). Preferred nucleobase sequences of probes may be chosen within positions 1306 to 1322 and 1621 to 1631 of the alignment corresponding to positions 2271 to 2285 and 2581 to 2591 of M. tuberculosis sequence (indicated by heavy frames). Mismatches between the sequences of M. avium, M. phlei, M. leprae, M. paratuberculosis, M. gastri and M. kansasii and that of M. tuberculosis in the alignment are indicated by light frames.

Figure 4 shows positions 2361 to 2520 and 3081 to 3120 of an alignment of 23S rRNA of M. tuberculosis (positions 3307 to 3466 and 4026 to 4064 of GenBank entry GB:MTCY130, accession number Z73992), M. avium (positions 2607 to 2765 and 3325 to 3363 of GenBank entry GB:MA23SRNA, accession number X74494), M. phlei (positions 2699 to 2858 ans 3418 to to 3456 of GenBank entry GB:MP23SRNA, accession number X74493), M. leprae (positions 2630 to 2789 of GenBank entry GB:ML5S23S, accession number X56657), M. paratuberculosis (positions 2607 to 2765 and 3325 to 3363 of GenBank entry GB:MPARRNA, accession number X74495), and M. kansasii (positions 2334 to 2494 and 3053 to 3091 of GenBank entry GB:MK23SRRNA, accession number Z17212). Preferred nucleobase sequences of probes may be chosen within positions 2401 to 2418, 2455 to 2486 and 3094 to 3103 of the alignment corresponding to positions 3347 to 3364, 3401 to 3432 and 4038 to 4047 of M. tuberculosis sequence (indicated by heavy frames). Mismatches between the sequences of M. avium, M. phlei, M. leprae, M. paratuberculosis, M. gastri and M. kansasii and that of M. tuberculosis in the alignment are indicated by light frames.

SPECIFIC DESCRIPTION

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The present invention provides novel probes for use in rapid and sensitive hybridisation based assays for the detection of organisms belonging to the MTC group.

We have identified suitable variable regions of the target nucleic acid by comparative analysis of generally available 23S rRNA sequences. Computers and computer programs which have been used for the purposes herein disclosed are generally available. In the probe design, sequence variations between the organisms belonging to the MTC group and other organisms have been taken into consideration, in particular M. avium.

When designing the probes, due regard should be taken to the assay conditions under which the probes are to be used. The stringency of the assay conditions determines the degree of complementarity needed between the probe and nucleic acid forming a hybrid. Stringency is chosen so as to maximise the difference in stability between the hybrid formed with the target nucleic acid and the non-target nucleic acid. It is desirable to have probes which hybridise under conditions of high stringency. Under such conditions, only highly complementary nucleic

acids will form stable hybrids with the probe according to the invention; stable hybrids without a sufficient degree of complementarity will not be formed.

Furthermore, probes should be positioned so as to minimise the stability of the probe:non-target nucleic acid hybrid. This may be accomplished by minimising the degree of complementarity to non-target nucleic acid and by designing the probe to span as many destabilising mismatches as possible. Whether a probe is useful to detect an organism belonging to the MTC group depends largely on the thermal stability difference between probe:target hybrids and probe:non-target hybrids. In designing the probes, the differences in these T_m values should be as large as possible.

Hybrids formed between peptide nucleic acid probes and nucleic acids have a higher thermal instability of mismatching bases compared to nucleic acid duplexes of the same sequences. Thus, the peptide nucleic acid probes exhibit a greater specificity for a complementary nucleic acid sequence than the traditional nucleic acid probe, which is seen as a greater difference in T_m values for probe:target hybrids and probe:non-target hybrids.

The length of the probe sequence is also important. The optimal length of a probe comprising a particular site of differences in base composition, e.g. among homologous regions of mycobacteria 23S rRNA, is a compromise between the principle that longer probes ensure specificity and shorter probes ensure that the destabilising differences in base composition constitute a greater portion of the probe.

Peptide nucleic acids can form duplexes in either orientation, but the antiparallel orientation form the most regular and stable duplex. Hence the antiparallel configuration is preferred for probe applications.

Mainly because the peptide nucleic acid strand is uncharged, a peptide nucleic acid-nucleic acid-duplex will have a higher T_m than the corresponding nucleic acid-nucleic acid-duplex. Typically there will be an increase in T_m of about 1 °C per base pair at 100 mM NaCl depending on the sequence (Egholm et al. (1993), Nature, 365, 566-568).

In contrast to DNA-DNA-duplex formation, no salt is necessary to facilitate and stabilise the formation of a peptide nucleic acid-DNA or a peptide nucleic acid-RNA duplex. The T_m of the peptide nucleic acid-DNA-duplex changes only a little with increasing ionic strength. Typically for a 15-mer, the T_m will drop only 5 °C when the salt concentration is raised from 10 mM NaCl to 1 M NaCl. At low ionic strength (e.g. 10 mM phosphate buffer with no salt added), it is possible to hybridise peptide nucleic acid to a target sequence under conditions where no

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stable DNA-DNA-duplex formation is able to occur (Nucleic Acid Hybridisation, a practical approach, eds. B. D. Hames & S. J. Higgins, IRL Press 1985, page 62-64). Furthermore, target sites that normally are inaccessible can be made more readily accessible for hybridisation with peptide nucleic acid probes at low salt concentration as the secondary and tertiary structure of nucleic acids are melted under such conditions.

Although it is preferred to use peptide nucleic acid probes targeting specific sequences of rRNA, it will readily be understood that peptide nucleic acid probes complementary to the rRNA targeting probes will be useful for the detection of the genes (DNA) coding for said sequence specific rRNA. Thus, as used herein, "probes able to form hybrids with target sequences in 23S rRNA" refers to probes capable of hybridising to sequences in 23S rRNA or to corresponding sequences in the non-coding strand of the rDNA as well as it refers to complementary probes capable of hybridising to the coding strand of DNA coding for the target rRNA sequences.

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In accordance with the present invention, peptide nucleic acid probes of formula (I) are provided, which probes are useful for detecting mycobacteria of the Mycobacterium tuberculosis Complex (MTC) in a sample, and which probes comprise from 10 to 30 polymerised moieties of formula (I)

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wherein each X and Y independently designate O or S, each Z independently designates O, S, NR^1 , or $C(R^1)_2$, wherein each R^1 independently designate H, C_{1-6} alkyl, C_{1-6} alkynyl, each R^2 , R^3 and R^4 independently designate H, the side chain of a naturally occurring amino acid, the side chain of a non-naturally occurring nucleobase, C_{1-4} alkyl, C_{1-4} alkenyl or C_{1-4} alkynyl, or a functional group, each Q independently designates a naturally occurring

nucleobase, a non-naturally occurring nucleobase, an intercalator, a nucleobase-binding

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with the proviso that the Qs of adjacent moieties are selected so as to form a sequence of which a subsequence includes at least one nucleobase complementary to a nucleobase of M. tuberculosis 23S rRNA that differs from the corresponding nucleobase of M. avium located

group, a label or H,

within the following domains

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Position 326 to Position 359 in Figure 1, or Position 635 to Position 658 in Figure 1, or Position 761 to Position 787 in Figure 2, or Position 1306 to Position 1322 in Figure 3, or Position 1621 to Position 1631 in Figure 3, or Position 2401 to Position 2418 in Figure 4, or Position 2455 to Position 2486 in Figure 4, or

Position 3094 to Position 3103 in Figure 4,

and further with the proviso that the probe comprising such subsequence is able to form hybrids with target sequences in 23S rRNA of said mycobacteria.

The term "naturally occurring nucleobases" includes the four main DNA bases (i.e. thymine (T), cytosine (C), adenine (A) and guanine (G)) as well as other naturally occurring nucleobases (e.g. uracil (U) and hypoxanthine).

The term "non-naturally occurring nucleobases" comprises i.a. modified naturally occurring nucleobases. Such non-naturally occurring nucleobases may be modified by substitution by e.g. one or more C₁₋₈ alkyl, C₁₋₈ alkenyl or C₁₋₈ alkynyl groups or labels. Examples of non-naturally occurring nucleobases are purine, 2,6-diamino purine, 5-propynylcytosine (C propynyl), isocytosine (iso-C), 5-methyl-isocytosine (iso^{Me}C) (see e.g. Tetrahedron Letters Vol 36, No 12, 2033-2036 (1995) or Tetrahedron Letters Vol 36, No 21, 3601-3604 (1995)), 7-deazaadenine, 7-deazaguanine, N⁴-ethanocytosine, N⁶-ethano-2,6-diaminopurine, 5-(C₃₋₆)-alkynylcytosine, 5-fluorouracil and pseudocytosine.

Examples of useful intercalators are e.g. acridin, antraquinone, psoralen and pyrene.

Examples of useful nucleobase-binding groups are e.g. groups containing cyclic or heterocyclic rings. Non-limiting examples are 3-nitro pyrrole and 5-nitro indole.

It is to be understood that alkyl, alkenyl and alkynyl groups may be branched or non-branched, cyclic or non-cyclic, and may further be interrupted by one or more heteroatoms, or may be unsubtituted or substituted by or may contain one or more functional groups. Non-limiting examples of such functional groups are acetyl groups, acyl groups, amino groups, carbamido groups, carbamyl groups, carbamyl groups, carbonyl groups, carboxy groups, cyano groups, dithio groups, formyl groups, guanidino groups, halogens, hydrazino groups, hydrazo groups,

hydroxamino groups, hydroxy groups, keto groups, mercapto groups, nitro groups, phospho groups, phospho ester groups, sulfo groups, thiocyanato groups, cyclic, aromatic and heterocyclic groups.

C₁₋₄ groups contain from 1 to 4 carbon atoms, C₁₋₆ groups contain from 1 to 6 carbon atoms, and C₁₋₁₅ groups contain from 1 to 15 carbon atoms, not including optional substituents, heteroatoms and/or functional groups. Non-limiting examples of such groups are -OH, -CH₃, -CF₃, -CH₂-, -CH₂CH₃, -CH₂CH₂-, -CH(CH₃)₂, -OCH₃, -OCH₂-, -OCH₂CH₃, -OCH₂CH₂-, -OCH(CH₃)₂, -OC(O)CH₃, -OC(O)CH₂-, -C(O)H, -C(O)-, -C(O)CH₃, -C(O)OH, -C(O)O-, -CH₂NH₂, -CH₂NH-, -CH₂OCH₃, -CH₂OCH₂-, -CH₂OC(O)OH, -CH₂OC(O)O-, -CH₂C(O)CH₃, -CH₂C(O)CH₂-, -C(O)NH₂, -P(O)₄H, -SH, -NH₂, -CH=CH₂, -CH=CH-, -CH=CHCH₂C(O)OH, -CH=CHCH₂C(O)O-, -C=CH, -C=C-, -CH₂C=CH, -CH₂C=C-, -CH₂C=CCH₃, -OCH₂C=CH, -OCH₂C=CO-, -OCH₂C=CCH₃, -NHCH₂C(O)-, -NHCH₂CH₂C(O)-, -NH(CH₂CH₂O)₂CH₂C(O)-, and HO(O)CCH₂C(O)(NH-(CH₂CH₂O)₂CH₂C(O))₂-, phenyl, benzyl, naphthyl, oxazolyl, pyridinyl, thiadiazolyl, triazolyl, and thienyl.

Within the present context, the expression "naturally occurring amino acid" is intended to comprise D- and L-forms of amino acids commonly found in nature, e.g. D- and L-forms of Ala (alanine), Arg (arginine), Asn (aspargine), Asp (aspartic acid), Cys (cysteine), Gln (glutamine), Glu (glutamic acid), His (histidine), Ile (isoleucine), Leu (leucine), Lys (lysine), Met (methionine), Phe (phenylalanine), Pro (proline), Ser (serine), Thr (threonine), Trp (tryptophan), Tyr (tyrosine) and Val (valine).

In the present context, the expression "non-naturally occurring amino acid" is intended to comprise D- and L-forms of amino acids other than those commonly found in nature as well as modified naturally occurring amino acids. Examples of useful non-naturally occurring amino acids are D- and L-forms of Cha (cyclohexylalanine), Cit (citrulline), Hci (homocitrulline), HomoCys (homocystein), Hse (homoserine), Nle (norleucine), Nva (norvaline), Orn (ornithine), Sar (sarcosine) and Thi (thienylalanine).

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The strength of the binding between the probe and the nucleic acid sequence is further influenced by the ligand Q. When Q designates a nucleobase, Hoogsteen and/or Watson-Crick base pairing assists in the formation of hybrids between a nucleic acid sequence to be detected and the probe. It is contemplated that one or more of the ligands may be a group which contribute little or none to the binding of the nucleic acid such as hydrogen. It is contemplated that suitable probes to be used comprise less than 25% by weight of moieties, wherein Q designates such groups. One or more of the ligands Q may be groups that stabilise nucleobase stacking such as intercalators or nucleobase-binding groups.

In the above-indicated probes one or more of the Q-groups may designate a label. Examples of suitable labels are given below. Moieties wherein Q denotes a label may preferably be located in one or both of the terminating moieties of the probe. Moieties wherein Q denotes a label may also be located internally.

The peptide nucleic acid probes may comprise moieties, wherein all X groups are O (polyamides) or wherein all X groups are S (polythioamides). It is to be understood that the probes may also comprise mixed moieties (comprising both amide and thioamide moieties).

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In another aspect, the present invention relates to peptide nucleic acid probes of formula (II), (III) and (IV)

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$$\sum_{\mathbb{R}^3} \mathbb{N}$$

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$$\sum_{\mathbb{R}^4} (|V|)$$

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wherein Z, R², R³, and R⁴, and Q is as defined above, which probes are suitable for detecting mycobacteria of the MTC.

In a preferred embodiment, the peptide nucleic acid probes according to the invention are of formulas (I)-(IV) as defined above with Z being NH, NCH₃ or O, each R^2 , R^3 and R^4 independently being the side chain of a naturally occurring nucleobase, the side chain of a non-naturally occurring nucleobase, or C_{1-4} alkyl, and each Q being a naturally occurring nucleobase or a non-naturally occurring nucleobase.

Peptide nucleic acid probes according to the invention are preferably those of formula (I)-(IV)

as defined above, wherein Z is NH or O, and R² is H or the side chain of Ala, Asp, Cys, Glu, His, HomoCys, Lys, Orn, Ser or Thr, and Q is a nucleobase selected from thymine, adenine, cytosine, guanine, uracil, iso-C, iso-G and 2,6-diaminopurine.

Peptide nucleic acid probes, which are of major interest for detecting mycobacteria of the MTC group, are probes of formula (V)

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wherein R^4 is H or the side chain of Ala, Asp, Cys, Glu, His, HomoCys, Lys, Orn, Ser or Thr, and Q is as defined above and with the provisos indicated above.

The peptide nucleic acid probe comprises polymerised moieties as defined above. From the formula, it is to be understood that the probe may comprise polymerised moieties which structure may be mutually different or identical. It may be advantageous that at least one moiety of the probe, preferably one (or both) of the moieties terminating the probe, are of a different structure. Such terminating moieties may suitably be a moiety of formula (VI)

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where Q is as defined above. Such moiety may suitably be connected to a peptide nucleic acid moiety though an amide bond.

The preferred length of the probe will depend on the target material and whether labelled probes are used. It is contemplated that especially interesting probes comprise from 10 to 30 polymerised moieties as defined above. Probes of the invention may suitably comprise from 12 to 25 polymerised moieties, more suitably from 14 to 22 polymerised moieties, most suitably from 15 to 20 polymerised moieties.

As mentioned above, the polymerised moieties of the probes may be mutually different or identical. In some embodiments, the polymerised moieties of formulas (V) constitute at least 75% by weight (calculated by excluding labels and linkers), preferably at least 80% by weight and most preferably at least 90% by weight of the probe.

The ends on the moieties terminating the probe may be substituted by suitable substituents which in the following will be named "linkers". A terminating end may suitably be substituted by from 1 to 5 linkers, more suitably from 1 to 3 linkers. Such linkers may suitably be selected among C_{1-15} alkyl, C_{1-15} alkenyl and C_{1-15} alkynyl groups as defined above. The linkers may be substituted or unsubstituted, branched or non-branched, or be interrupted by heteroatoms, or be substituted or contain functional groups as described above. This may depend on the chemical nature of the terminating moiety (i.e. whether the moiety is terminated by a carbon, oxygen or nitrogen atom). A terminating end or a linker on a terminating end may further be substituted by one or more labels, which labels may be incorporated end to end, i.e. so as to form a non-branched labelled end, or may be incorporated so as to form a branched labelled end ("zipper"). The linkers may be attached directly to a terminating end, may be attached to a label or between labels on a terminating end, or be attached to a terminating end before a label is attached to a terminating end. It should be understood that two terminating ends may carry different or identical substituents, linkers and/or labels. It should further be understood that the term "a label" is intended to comprise one or more labels as the term "linkers" is to comprise one or more linkers.

Examples of suitable linkers are -NH(CH $_2$ CH $_2$ O) $_n$ CH $_2$ C(O)-, -NH(CHOH) $_n$ C(O)-, -(O)C(CH $_2$ OCH $_2$) $_n$ C(O)- and -NH(CH $_2$) $_n$ C(O)-, NH $_2$ (CH $_2$ CH $_2$ O) $_n$ CH $_2$ C(O)-, NH $_2$ (CHOH) $_n$ C(O)-, HO(O)C(CH $_2$ OCH $_2$) $_n$ C(O)-, NH $_2$ (CH $_2$) $_n$ C(O)-, -NH(CH $_2$ CH $_2$ O) $_n$ CH $_2$ C(O)OH, -NH(CHOH) $_n$ C(O)OH, -(O)C(CH $_2$ OCH $_2$) $_n$ C(O)OH and -NH(CH $_2$) $_n$ C(O)OH, wherein n is 0 or an integer from 1 to 8, preferably from 1 to 3. Examples of very interesting linkers are -NHCH $_2$ C(O)-, -NHCH $_2$ CH $_2$ C(O)-, -NH(CH $_2$ CH $_2$ O) $_2$ CH $_2$ C(O)-, HO(O)CCH $_2$ CH(O)(NH-(CH $_2$ CH $_2$ O) $_2$ CH $_2$ C(O)) $_2$ -.

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In the present context, the term "label" refers to a substituent which is useful for detection or visualisation. Suitable labels comprise fluorophores, biotin, dinitro benzoic acid, digoxigenin, radioisotope labels, peptide or enzyme labels, chemiluminiscence labels, hapten, antigen or antibody labels.

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The expression "peptide label" is intended to mean a label comprising from 1 to 20 naturally occurring or non-naturally occurring amino acids, preferably from 1 to 10 naturally occurring or non-naturally occurring amino acids, more preferably from 1 to 8 naturally occurring or non-naturally occurring amino acids, most preferably from 1 to 4 naturally occurring or non-naturally occurring amino acids, linked together end to end in a non-branched or branched ("zipper") fashion. In a preferred embodiment, such a non-branched or branched end comprises one or more, preferably from 1 to 8 labels, more preferably from 1 to 4, further labels other than a peptide label. Such further labels may suitably terminate a non-branched

end or a branched end. One or more linkers may suitably be attached to the terminating end before a peptide label and/or a further label is attached. Such linker units may also be attached between a peptide label and a further label.

- The probe as such may also comprise one or more labels such as from 1 to 8, preferably from 1 to 4, labels and/or one or more linker units, which may be attached internally, i.e. to the backbone of the probe. The linker units and labels may mutually be attached as described above.
- Examples of particular interesting labels are biotin, fluorescent labels, such as fluorescein labels, e.g. 5-(and 6)-carboxyfluorescein, 5- or 6-carboxyfluorescein, 6-(fluorescein)-5-(and 6)-carboxamido hexanoic acid and fluorescein isothiocyanate, peptide labels consisting of from 1 to 20 naturally occurring amino acids or non-naturally occurring amino acids, peroxidases such as horse radish peroxidase (HRP) and soya bean peroxidase, dinitro benzoic acid, rhodamine, tetramethylrhodamine, cyanine dyes such as Cy2, Cy3 and Cy5, coumarin, R-phycoerythrin (RPE), allophycoerythrin, Texas Red and Princeton Red as well as conjugates of R-phycoerythrin and, e.g. Cy5 or Texas Red.

Examples of preferred labels are biotin, fluorescent labels, peptide labels and dinitro benzoic acid. Peptide labels may preferably be composed of from 1 to 10, more preferably of from 1 to 8, most preferably of from 1 to 4, naturally occurring or non-naturally occurring amino acids. It may be particularly advantageous to incorporate one or more labels other as well as a peptide label such as from 1 to 8 or from 1 to 4 other labels.

25 Suitable peptide labels may preferably be composed of cysteine, glycine, lysine or ornithine.

In a further embodiment, the Q substituent as defined above may be labelled. Suitable labels are as defined above. Between Q and such a label, a linker as defined above may be incorporated. It is preferred that such labelled ligands Q are selected from thymine and uridine labelled in the 5-position and 7-deazaguanine and 7-deazaguanine labelled in the 7-position.

The probes may be synthesised according to the procedures described in "PNA Information Package" obtained from Millipore Corporation (Bedford, MA, USA), or may be synthesised on an Expedite Nucleic Acid Synthesis System (PerSeptive, USA).

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If using the Fmoc strategy for elongation of the probe with linkers or amino acids, it was possible to retain side chain amino groups protected with acid sensitive protection groups such as the Boc or Mtt group. This method allows introduction of a linker containing several

Boc protected amino groups which can all be cleaved and labelled in the same synthesis cycle.

One way of labelling a probe is to use a fluorescent label, such as 5-(and 6)-carboxyfluorescein, 5- or 6-carboxyfluorescein, or 6-(fluorescein)-5-(and 6)-carboxamido hexanoic acid. The acid group is activated with HATU (O-(7-azabenzotriazol-1-yl)-1,1,3,3-tetramethyluronium hexafluorophosphate) or HBTU (2-(1H-benzotriazol-1-yl)-1,1,3,3-tetramethyluronium hexafluorophosphate) and reacted with the N-terminal amino group. The same technique can be applied to other labelling groups containing an acid function. Alternatively, the succinimidyl ester of the above-mentioned labels or fluorescein isothiocyanate may be used directly.

After synthesis, probes were cleaved from the resin using standard procedures as described by Millipore Corporation or PerSeptive Biosystems. The probes were purified and analysed using reversed-phase HPLC techniques at 50°C and were characterised by matrix-assisted laser desorption/ionisation time of flight mass spectrometry (MALDI-TOFMS), plasma desorption mass spectrometry (PDMS), electron spray mass spectrometry (ESMS), or fast atom bombardment (FAB-MS).

Generally, probes such as probes comprising polymerised moieties of formula (V) may also be prepared as described in, e.g., Tetrahedron Letters Vol 35, No 29, 5173-5176 (1994) and Bioorganic & Medical Chemistry Letters, Vol 4, No 8, 1077-1080 (1994). Chemical properties of some probes are described in, e.g., Nature, 365, 566-568 (1993).

Detection of the label depend on the type of label and on the format of the procedure. In cases where the sample is deposited onto slides, the hybridisation results may be visualised using well known immunohistochemical staining methods to detect the labelling on the probe. When fluorescent labelled binding partners are used, the hybrids may be detected using an antibody against the fluorescent label which antibody may be conjugated with an enzyme. The fluorescent label may alternatively be detected directly using a fluorescence microscope, or the results may be automatically analysed on a fluorescent-based image analysis system.

When biotin labelled probes are used, the hybrids may be detected using an antibody against the biotin label which antibody may be conjugated with an enzyme. If necessary, an enhancement of the signal can be generated using commercially available amplification systems such as the catalysed signal amplification system for biotinylated probes (DAKO K 1500).

The probes according to the invention are used in the detection of mycobacteria of the MTC in

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samples which may contain these bacteria.

In the assay method, at least one probe according to the invention is contacted with target nucleic acid and an analysis for hybrid formation is carried out.

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In the assay method according to the invention, a sample to be analysed for the presence of mycobacteria of the MTC is contacted with one or more probes according to the invention under such conditions by which hybridisation between the probe and any complementary sample rRNA of mycobacteria of the MTC can occur, and observing or measuring the resulting hybridisation.

In one embodiment of the assay method, conventionally prepared smears of bacterial cells are contacted with one or more probes according to the invention under conditions suitable for hybridisation to occur between the probe(s) and any complementary rRNA in the sample. The complexes formed are detected. An example of this assay format is fluorescence *in situ* hybridisation (FISH), wherein the probes according to the invention are labelled with fluorescein or another fluorophore. When designing MTC probes, it might be advantageous to use more than one probe. If e.g. three such probes are included in the assay each in a concentration of one third of the concentration of a single probe, possible cross reactivity of the individual probes will not invalidate the results.

In another embodiment of the assay method, a test sample is firstly subjected to conditions, which release nucleic acid from the bacteria present in that sample. Contact between one or more probes as defined herein, which may be labelled, and the rRNA target may be carried out in solution under conditions, which promote hybridisation between the probe(s) and any target nucleic acid present. The probe:nucleic acid complex may be immobilised to a solid support, e.g. by using a capture probe.

Due to the high affinity of the probes defined herein for nucleic acids, it is not necessary to carry out the hybridisation of the probe and nucleic acid in solution. This allows flexibility in the assay format. For instance, the detection probes can be brought into contact with the target nucleic acid in solution and the probe/nucleic acid complex can be captured by an immobilised capture probe. Or the sample comprising the target nucleic acid can even be added to an assay system comprising detection probes as well as immobilised capture probe. The immobilisation of the capture probe may be effected by using a streptavidin coated solid phase and a biotinylated capture probe. The probe may be immobilised onto a solid support by coupling reaction between a carboxylic acid on the linker and an amino derivatised support. Alternatively, the coupling onto the solid support may be accomplished by photochemical

activation of photoreactive groups which have been attached absorptively to the solid support prior to photochemical activation. Such photoreactive groups are described in EP 408 078 A.

In practice, a solid phase based assay system is very attractive as the analysis can be carried out using a solid phase precoated with a capture probe. A solid phase based assay system is also feasible for automatisation of the analysis.

The capture probe may be one of the other MTC probes not used in the hybridisation reaction and detection step for target nucleic acid, thus ensuring dual species specificity. The dual specificity will allow shorter probes be used, e.g. 10 mer probes.

The solid support capture system may take a wide variety of forms well known in the art, such as e.g. a plate, a microtiter plate having one or more wells, a microscope slide, a filter, a membrane, a tube, a dip stick, a strip, beads such as paramagnetic beads, beads made of polystyrene, polypropylene, polyethylene, dextran, nylon, amyloses, natural and modified celluloses, polyacrylamides and agaroses. When a filter, a membrane, a strip or beads is (are) used as the solid support, it (they) may, if conveniently be incorporated into a single-use.

It has been observed that peptide nucleic acids may bind to a variety of solid phases. A blocking reaction is required to reduce non-specific binding of the peptide nucleic acids to the solid phase. The blocking reaction may be carried out with commonly used blocking reagents, such as BSA (bovine serum albumin), casein, Triton X-100® or Tween 20®. The preferred blocking reagents are Triton X-100® and Tween 20®.

The captured probe:nucleic acid complexes may be detected or identified by a wide variety of methods for that purpose. The probe to be brought in contact with the target nucleic acid may be labelled, whereby said may form part of the detection system. In another embodiment, the captures probe:nucleic acid complexes are detected using a detection system based on an antibody reacting specifically with complexes formed between peptide nucleic acid and nucleic acid (such as described in WO 95/17430), in which detection system the primary antibody may comprise a label, or which detection system comprises a labelled secondary antibody, which specifically binds to the primary antibody.

DESCRIPTION OF SPECIFIC EMBODIMENTS

Examples of suitable Qs of adjacent moieties are given below. Peptide nucleic acid probes comprising such Qs will be able to detect mycobacteria of the MTC group. The probes are written from left to right corresponding to from the C-terminal end towards the N-terminal end.

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The nucleobase mismatche(s) between M. tuberculosis and M. avium is (are) indicated in bold.

Examples of nucleobase sequences of peptide nucleic acid probes wherein the Qs of adjacent
moieties are selected so as to be complementary to nucleobases within Position 326 to 359
shown in Figure 1 are the following
AGC GCT GAG ACA TAT CCT CCC A
GC GCT GAG ACA TAT CCT CCC A
C GCT GAG ACA TAT CCT CCC A

10 AGC GCT GAG ACA TAT CCT CCC
AGC GCT GAG ACA TAT CCT CC
AGC GCT GAG ACA TAT CCT CC
AGC GCT GAG ACA TAT CCT CC
AGC GCT GAG ACA TAT CCT C

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TAG CGC TGA GAC ATA TCC TCC C
AG CGC TGA GAC ATA TCC TCC C
G CGC TGA GAC ATA TCC TCC C
CGC TGA GAC ATA TCC TCC C

TAG CGC TGA GAC ATA TCC TCC
TAG CGC TGA GAC ATA TCC TC
TAG CGC TGA GAC ATA TCC TC
TAG CGC TGA GAC ATA TCC T

25 GTA GCG CTG AGA CAT ATC CTC C
TA GCG CTG AGA CAT ATC CTC C
A GCG CTG AGA CAT ATC CTC C
GCG CTG AGA CAT ATC CTC C
CG CTG AGA CAT ATC CTC C
30 GTA GCG CTG AGA CAT ATC CTC
GTA GCG CTG AGA CAT ATC CT
GTA GCG CTG AGA CAT ATC CT

GGT AGC GCT GAG ACA TAT CCT C

35 GT AGC GCT GAG ACA TAT CCT C

T AGC GCT GAG ACA TAT CCT C

AGC GCT GAG ACA TAT CCT C

GC GCT GAG ACA TAT CCT C

C GCT GAG ACA TAT CCT C
GGT AGC GCT GAG ACA TAT CCT
GGT AGC GCT GAG ACA TAT CC

5 GGG TAG CGC TGA GAC ATA TCC T
GG TAG CGC TGA GAC ATA TCC T
G TAG CGC TGA GAC ATA TCC T
TAG CGC TGA GAC ATA TCC T
AG CGC TGA GAC ATA TCC T

10 G CGC TGA GAC ATA TCC T
CGC TGA GAC ATA TCC T
GGG TAG CGC TGA GAC ATA TCC

CGG GTA GCG CTG AGA CAT ATC C

GG GTA GCG CTG AGA CAT ATC C

G GTA GCG CTG AGA CAT ATC C

GTA GCG CTG AGA CAT ATC C

TA GCG CTG AGA CAT ATC C

A GCG CTG AGA CAT ATC C

20 GCG CTG AGA CAT ATC C

CG CTG AGA CAT ATC C

CCG GGT AGC GCT GAG ACA TAT C
CG GGT AGC GCT GAG ACA TAT C
25 G GGT AGC GCT GAG ACA TAT C
GGT AGC GCT GAG ACA TAT C
GT AGC GCT GAG ACA TAT C
T AGC GCT GAG ACA TAT C
AGC GCT GAG ACA TAT C
CGCT GAG ACA TAT C
CGCT GAG ACA TAT C

GCC GGG TAG CGC TGA GAC ATA T
CC GGG TAG CGC TGA GAC ATA T

35 C GGG TAG CGC TGA GAC ATA T
GGG TAG CGC TGA GAC ATA T
GCC GGG TAG CGC TGA GAC ATA
GCC GGG TAG CGC TGA GAC ATA

GCC GGG TAG CGC TGA GAC A
GCC GGG TAG CGC TGA GAC
GCC GGG TAG CGC TGA GA
GCC GGG TAG CGC TGA G
GCC GGG TAG CGC TGA
GCC GGG TAG CGC TGA

AGC CGG GTA GCG CTG AGA CAT A GC CGG GTA GCG CTG AGA CAT A C CGG GTA GCG CTG AGA CAT A 10 CGG GTA GCG CTG AGA CAT A GG GTA GCG CTG AGA CAT A AGC CGG GTA GCG CTG AGA CAT AGC CGG GTA GCG CTG AGA CA AGC CGG GTA GCG CTG AGA C 15 AGC CGG GTA GCG CTG AGA AGC CGG GTA GCG CTG AG AGC CGG GTA GCG CTG A AGC CGG GTA GCG CTG AGC CGG GTA GCG CT 20

CAG CCG GGT AGC GCT GAG ACA T AG CCG GGT AGC GCT GAG ACA T G CCG GGT AGC GCT GAG ACA T CCG GGT AGC GCT GAG ACA T 25 CG GGT AGC GCT GAG ACA T G GGT AGC GCT GAG ACA T CAG CCG GGT AGC GCT GAG ACA CAG CCG GGT AGC GCT GAG AC CAG CCG GGT AGC GCT GAG A 30 CAG CCG GGT AGC GCT GAG CAG CCG GGT AGC GCT GA CAG CCG GGT AGC GCT G CAG CCG GGT AGC GCT CAG CCG GGT AGC GC 35

> CAG CCG GGT AGC GCT GAG ACA T AG CCG GGT AGC GCT GAG ACA T

G CCG GGT AGC GCT GAG ACA T
CCG GGT AGC GCT GAG ACA T
CG GGT AGC GCT GAG ACA T
CAG CCG GGT AGC GCT GAG ACA
5 CAG CCG GGT AGC GCT GAG AC
CAG CCG GGT AGC GCT GAG A
CAG CCG GGT AGC GCT GAG
CAG CCG GGT AGC GCT GA
CAG CCG GGT AGC GCT GA
CAG CCG GGT AGC GCT GA
CAG CCG GGT AGC GCT G
10 CAG CCG GGT AGC GCT
CAG CCG GGT AGC GCT

TCA GCC GGG TAG CGC TGA GAC A CA GCC GGG TAG CGC TGA GAC A A GCC GGG TAG CGC TGA GAC A 15 GCC GGG TAG CGC TGA GAC A CC GGG TAG CGC TGA GAC A C GGG TAG CGC TGA GAC A GGG TAG CGC TGA GAC A TCA GCC GGG TAG CGC TGA GAC 20 TCA GCC GGG TAG CGC TGA GA TCA GCC GGG TAG CGC TGA G TCA GCC GGG TAG CGC TGA TCA GCC GGG TAG CGC TG TCA GCC GGG TAG CGC T 25 TCA GCC GGG TAG CGC TCA GCC GGG TAG CG

CTC AGC CGG GTA GCG CTG AGA C

TC AGC CGG GTA GCG CTG AGA C

C AGC CGG GTA GCG CTG AGA C

AGC CGG GTA GCG CTG AGA C

GC CGG GTA GCG CTG AGA C

C CGG GTA GCG CTG AGA C

C CGG GTA GCG CTG AGA C

GG GTA GCG CTG AGA C

CTC AGC CGG GTA GCG CTG AGA

CTC AGC CGG GTA GCG CTG AGA

CTC AGC CGG GTA GCG CTG A
CTC AGC CGG GTA GCG CTG
CTC AGC CGG GTA GCG CT
CTC AGC CGG GTA GCG C

CTC AGC CGG GTA GCG
CTC AGC CGG GTA GCG
CTC AGC CGG GTA GC

TCT CAG CCG GGT AGC GCT GAG A CT CAG CCG GGT AGC GCT GAG A T CAG CCG GGT AGC GCT GAG A 10 CAG CCG GGT AGC GCT GAG A AG CCG GGT AGC GCT GAG A G CCG GGT AGC GCT GAG A CCG GGT AGC GCT GAG A CG GGT AGC GCT GAG A 15 G GGT AGC GCT GAG A TCT CAG CCG GGT AGC GCT GAG TCT CAG CCG GGT AGC GCT GA TCT CAG CCG GGT AGC GCT G TCT CAG CCG GGT AGC GCT 20 TCT CAG CCG GGT AGC GC TCT CAG CCG GGT AGC G TCT CAG CCG GGT AGC

CTC TCA GCC GGG TAG CGC TGA G 25 TC TCA GCC GGG TAG CGC TGA G C TCA GCC GGG TAG CGC TGA G TCA GCC GGG TAG CGC TGA G CA GCC GGG TAG CGC TGA G A GCC GGG TAG CGC TGA G 30 GCC GGG TAG CGC TGA G CC GGG TAG CGC TGA G C GGG TAG CGC TGA G GGG TAG CGC TGA G CTC TCA GCC GGG TAG CGC TGA 35 CTC TCA GCC GGG TAG CGC TG CTC TCA GCC GGG TAG CGC T CTC TCA GCC GGG TAG CGC

CTC TCA GCC GGG TAG CG

CCT CTC AGC CGG GTA GCG CTG A

T CTC AGC CGG GTA GCG CTG A

T CTC AGC CGG GTA GCG CTG A

CTC AGC CGG GTA GCG CTG A

CTC AGC CGG GTA GCG CTG A

TC AGC CGG GTA GCG CTG A

C AGC CGG GTA GCG CTG A

GC CGG GTA GCG CTG A

GC CGG GTA GCG CTG A

CCT CTC AGC CGG GTA GCG CTG

CCT CTC AGC CGG GTA GCG CT

15 CCT CTC AGC CGG GTA GCG C

CCT CTC AGC CGG GTA GCG C

CCT CTC AGC CGG GTA GCG

GCC TCT CAG CCG GGT AGC GCT G CC TCT CAG CCG GGT AGC GCT G 20 C TCT CAG CCG GGT AGC GCT G TCT CAG CCG GGT AGC GCT G CT CAG CCG GGT AGC GCT G T CAG CCG GGT AGC GCT G TCT CAG CCG GGT AGC GCT G 25 CT CAG CCG GGT AGC GCT G T CAG CCG GGT AGC GCT G CAG CCG GGT AGC GCT G AG CCG GGT AGC GCT G G CCG GGT AGC GCT G 30 GCC TCT CAG CCG GGT AGC GCT GCC TCT CAG CCG GGT AGC GC GCC TCT CAG CCG GGT AGC G GCC TCT CAG CCG GGT AGC GCC TCT CAG CCG GGT AG 35 GCC TCT CAG CCG GGT A GCC TCT CAG CCG GGT GCC TCT CAG CCG GG

- TGC CTC TCA GCC GGG TAG CGC TGC CTC TCA GCC GGG TAG CGC TCTC TCA GCC GGG TAG CGC TCTC TCA GCC GGG TAG CGC T
- TC TCA GCC GGG TAG CGC T
 TC TCA GCC GGG TAG CGC T
 C TCA GCC GGG TAG CGC T
 TCA GCC GGG TAG CGC T
 CA GCC GGG TAG CGC T
- 10 A GCC GGG TAG CGC T
 TGC CTC TCA GCC GGG TAG CGC
 TGC CTC TCA GCC GGG TAG CG
 TGC CTC TCA GCC GGG TAG C
 TGC CTC TCA GCC GGG TAG
- TGC CTC TCA GCC GGG TA
 TGC CTC TCA GCC GGG T
 TGC CTC TCA GCC GGG
 TGC CTC TCA GCC GG
- 20 CTG CCT CTC AGC CGG GTA GCG C
 TG CCT CTC AGC CGG GTA GCG C
 G CCT CTC AGC CGG GTA GCG C
 CCT CTC AGC CGG GTA GCG C
 CT CTC AGC CGG GTA GCG C
- 25 T CTC AGC CGG GTA GCG C
 CTC AGC CGG GTA GCG C
 TC AGC CGG GTA GCG C
 C AGC CGG GTA GCG C
 CTG CCT CTC AGC CGG GTA GCG
- CTG CCT CTC AGC CGG GTA GC
 CTG CCT CTC AGC CGG GTA G
 CTG CCT CTC AGC CGG GTA
 CTG CCT CTC AGC CGG GT
 CTG CCT CTC AGC CGG G

 35 CTG CCT CTC AGC CGG

ACT GCC TCT CAG CCG GGT AGC G

CTG CCT CTC AGC CG

CT GCC TCT CAG CCG GGT AGC G
T GCC TCT CAG CCG GGT AGC G
GCC TCT CAG CCG GGT AGC G
CC TCT CAG CCG GGT AGC G
CC TCT CAG CCG GGT AGC G
TCT CAG CCG GGT AGC G
TCT CAG CCG GGT AGC G
CT CAG CCG GGT AGC G
T CAG CCG GGT AGC G
ACT GCC TCT CAG CCG GGT AGC
ACT GCC TCT CAG CCG GGT A
ACT GCC TCT CAG CCG GGT A
ACT GCC TCT CAG CCG GGT
ACT GCC TCT CAG CCG GGT
ACT GCC TCT CAG CCG GGT
ACT GCC TCT CAG CCG GG

ACT GCC TCT CAG CCG

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GAC TGC CTC TCA GCC GGG TAG C AC TGC CTC TCA GCC GGG TAG C C TGC CTC TCA GCC GGG TAG C TGC CTC TCA GCC GGG TAG C 20 GC CTC TCA GCC GGG TAG C C CTC TCA GCC GGG TAG C CTC TCA GCC GGG TAG C TC TCA GCC GGG TAG C C TCA GCC GGG TAG C 25 GAC TGC CTC TCA GCC GGG TAG GAC TGC CTC TCA GCC GGG TA GAC TGC CTC TCA GCC GGG T GAC TGC CTC TCA GCC GGG GAC TGC CTC TCA GCC GG 30 GAC TGC CTC TCA GCC G GAC TGC CTC TCA GCC GAC TGC CTC TCA GC

Examples of nucleobase sequences of peptide nucleic acid probes wherein the Qs of adjacent moieties are selected so as to be complementary to nucleobases within Position 635 to 658 shown in Figure 1 are the following

GGA GAG GAA AAG GAG GCT CTG A

GGA GAG GAA AAG GAG GCT CTG
GGA GAG GAA AAG GAG GCT CT
GGA GAG GAA AAG GAG GCT C
GGA GAG GAA AAG GAG GCT

GGA GAG GAA AAG GAG GCT
GGA GAG GAA AAG GAG GC
GGA GAG GAA AAG GAG
GGA GAG GAA AAG GAG
GGA GAG GAA AAG GA
GGA GAG GAA AAG GA

10

CGG AGA GGA AAA GGA GGC TCT G
CGG AGA GGA AAA GGA GGC TCT
CGG AGA GGA AAA GGA GGC TC
CGG AGA GGA AAA GGA GGC T

15 CGG AGA GGA AAA GGA GGC
CGG AGA GGA AAA GGA GG
CGG AGA GGA AAA GGA G
CGG AGA GGA AAA GGA
CGG AGA GGA AAA GGA
CGG AGA GGA AAA GGA

20

CCG GAG AGG AAA AGG AGG CTC T
CCG GAG AGG AAA AGG AGG CTC
CCG GAG AGG AAA AGG AGG CT
CCG GAG AGG AAA AGG AGG C

25 CCG GAG AGG AAA AGG AGG
CCG GAG AGG AAA AGG AG
CCG GAG AGG AAA AGG A
CCG GAG AGG AAA AGG C
CCG GAG AGG AAA AGG
CCG GAG AGG AAA AGG

30

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TCC GGA GAG GAA AAG GAG GCT C
TCC GGA GAG GAA AAG GAG GCT
TCC GGA GAG GAA AAG GAG GC
TCC GGA GAG GAA AAG GAG G
TCC GGA GAG GAA AAG GAG
TCC GGA GAG GAA AAG GA
TCC GGA GAG GAA AAG G
TCC GGA GAG GAA AAG G

TCC GGA GAG GAA AA

CTC CGG AGA GGA AAA GGA GGC T
CTC CGG AGA GGA AAA GGA GGC

5 CTC CGG AGA GGA AAA GGA GG
CTC CGG AGA GGA AAA GGA G
CTC CGG AGA GGA AAA GGA
CTC CGG AGA GGA AAA GG
CTC CGG AGA GGA AAA G

CTC CGG AGA GGA AAA G

CTC CGG AGA GGA AAA
CTC CGG AGA GGA AAA

CCT CCG GAG AGG AAA AGG AGG C
CCT CCG GAG AGG AAA AGG AGG

15 CCT CCG GAG AGG AAA AGG AG
CCT CCG GAG AGG AAA AGG A
CCT CCG GAG AGG AAA AGG
CCT CCG GAG AGG AAA AG
CCT CCG GAG AGG AAA A

20 CCT CCG GAG AGG AAA
CCT CCG GAG AGG AAA

TCC TCC GGA GAG GAA AAG GAG G
TCC TCC GGA GAG GAA AAG GAG

25 TCC TCC GGA GAG GAA AAG GA
TCC TCC GGA GAG GAA AAG G
TCC TCC GGA GAG GAA AAG
TCC TCC GGA GAG GAA AA
TCC TCC GGA GAG GAA A

30 TCC TCC GGA GAG GAA
TCC TCC GGA GAG GAA

CTC CTC CGG AGA GGA AAA GGA G
CTC CTC CGG AGA GGA AAA GGA
35 CTC CTC CGG AGA GGA AAA GG
CTC CTC CGG AGA GGA AAA G
CTC CTC CGG AGA GGA AAA
CTC CTC CGG AGA GGA AAA

CTC CTC CGG AGA GGA A
CTC CTC CGG AGA GGA
CTC CTC CGG AGA GG

- 5 CCT CCT CCG GAG AGG AAA AGG A
 CCT CCT CCG GAG AGG AAA AGG
 CCT CCT CCG GAG AGG AAA AG
 CCT CCT CCG GAG AGG AAA A
 CCT CCT CCG GAG AGG AAA

 10 CCT CCT CCG GAG AGG AA
 CCT CCT CCG GAG AGG A
 CCT CCT CCG GAG AGG A
 CCT CCT CCG GAG AGG
 CCT CCT CCG GAG AGG
- 15 CCC TCC TCC GGA GAG GAA AAG G
 CCC TCC TCC GGA GAG GAA AAG
 CCC TCC TCC GGA GAG GAA AA
 CCC TCC TCC GGA GAG GAA A
 CCC TCC TCC GGA GAG GAA
 20 CCC TCC TCC GGA GAG GA
 CCC TCC TCC GGA GAG GA
 CCC TCC TCC GGA GAG
 CCC TCC TCC GGA GAG
 CCC TCC TCC GGA GAG
 CCC TCC TCC GGA GAG

25

ACC CTC CTC CGG AGA GGA AAA G
ACC CTC CTC CGG AGA GGA AAA
ACC CTC CTC CGG AGA GGA AA
ACC CTC CTC CGG AGA GGA A

30 ACC CTC CTC CGG AGA GGA
ACC CTC CTC CGG AGA GG
ACC CTC CTC CGG AGA G

ACC CTC CTC CGG AGA G

CC CTC CTC CGG AGA
CC CTC CTC CGG AGA GGA AAA G
CTC CTC CGG AGA GGA AAA G
CTC CTC CGG AGA GGA AAA G

C CTC CGG AGA GGA AAA G CTC CGG AGA GGA AAA G TC CTC CGG AGA GGA AAA G C CTC CGG AGA GGA AAA G

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CAC CCT CCT CCG GAG AGG AAA A CAC CCT CCT CCG GAG AGG AAA CAC CCT CCT CCG GAG AGG AA CAC CCT CCT CCG GAG AGG A 10 CAC CCT CCT CCG GAG AGG CAC CCT CCT CCG GAG AG CAC CCT CCT CCG GAG A CAC CCT CCT CCG GAG CAC CCT CCT CCG GA AC CCT CCT CCG GAG AGG AAA A

> TCA CCA CCC TCC TCC GG CA CCA CCC TCC TCC GG A CCA CCC TCC TCC GG CCA CCC TCC TCC GG TCA CCA CCC TCC TCC G TCA CCA CCC TCC TCC TCA CCA CCC TCC TC

Examples of nucleobase sequences of peptide nucleic acid probes wherein the Qs of adjacent 25 moieties are selected so as to be complementary to nucleobases within Position 761 to 787

shown in Figure 2 are the following ATG CGC GTG TGG GTC GC TG CGC GTG TGG GTC GC G CGC GTG TGG GTC GC 30 CGC GTG TGG GTC GC ATG CGC GTG TGG GTC G ATG CGC GTG TGG GTC ATG CGC GTG TGG GT

35

CGC GTA TGC GCG TGT GGG TCG C GC GTA TGC GCG TGT GGG TCG C C GTA TGC GCG TGT GGG TCG C

GTA TGC GCG TGT GGG TCG C

TA TGC GCG TGT GGG TCG C

A TGC GCG TGT GGG TCG C

TGC GCG TGT GGG TCG C

GC GCG TGT GGG TCG C

C GCG TGT GGG TCG C

CGC GTA TGC GCG TGT GGG TCG

CGC GTA TGC GCG TGT GGG TC

CGC GTA TGC GCG TGT GGG

CGC GTA TGC GCG TGT GGG

CGC GTA TGC GCG TGT GG

CGC GTA TGC GCG TGT GC

CGC GTA TGC GCG TGT G

CGC GTA TGC GCG TGT C

CGC GTA TGC GCG TGT C

15

GCG CGT ATG CGC GTG TGG GTC G CG CGT ATG CGC GTG TGG GTC G G CGT ATG CGC GTG TGG GTC G CGT ATG CGC GTG TGG GTC G GT ATG CGC GTG TGG GTC G 20 T ATG CGC GTG TGG GTC G ATG CGC GTG TGG GTC G TG CGC GTG TGG GTC G G CGC GTG TGG GTC G GCG CGT ATG CGC GTG TGG GTC 25 GCG CGT ATG CGC GTG TGG GT GCG CGT ATG CGC GTG TGG G GCG CGT ATG CGC GTG TGG GCG CGT ATG CGC GTG TG GCG CGT ATG CGC GTG T 30 GCG CGT ATG CGC GTG GCG CGT ATG CGC GT

CGC GCG TAT GCG CGT GTG GGT C

35 GC GCG TAT GCG CGT GTG GGT C

C GCG TAT GCG CGT GTG GGT C

GCG TAT GCG CGT GTG GGT C

CG TAT GCG CGT GTG GGT C

G TAT GCG CGT GTG GGT C

TAT GCG CGT GTG GGT C

AT GCG CGT GTG GGT C

T GCG CGT GTG GGT C

5 CGC GCG TAT GCG CGT GTG GGT

CGC GCG TAT GCG CGT GTG GG

CGC GCG TAT GCG CGT GTG

CGC GCG TAT GCG CGT GTG

CGC GCG TAT GCG CGT GT

10 CGC GCG TAT GCG CGT G

CGC GCG TAT GCG CGT C

CGC GCG TAT GCG CGT C

ACG CGC GTA TGC GCG TGT GGG T CG CGC GTA TGC GCG TGT GGG T 15 G CGC GTA TGC GCG TGT GGG T CGC GTA TGC GCG TGT GGG T GC GTA TGC GCG TGT GGG T C GTA TGC GCG TGT GGG T GTA TGC GCG TGT GGG T 20 TA TGC GCG TGT GGG T A TGC GCG TGT GGG T ACG CGC GTA TGC GCG TGT GGG ACG CGC GTA TGC GCG TGT GG ACG CGC GTA TGC GCG TGT G 25 ACG CGC GTA TGC GCG TGT ACG CGC GTA TGC GCG TG ACG CGC GTA TGC GCG T ACG CGC GTA TGC GCG ACG CGC GTA TGC GC 30

CAC GCG CGT ATG CGC GTG TGG G
AC GCG CGT ATG CGC GTG TGG G
C GCG CGT ATG CGC GTG TGG G
35 GCG CGT ATG CGC GTG TGG G
CG CGT ATG CGC GTG TGG G
G CGT ATG CGC GTG TGG G
CGT ATG CGC GTG TGG G

GT ATG CGC GTG TGG G

T ATG CGC GTG TGG G

CAC GCG CGT ATG CGC GTG TGG

CAC GCG CGT ATG CGC GTG TG

5 CAC GCG CGT ATG CGC GTG T

CAC GCG CGT ATG CGC GTG

CAC GCG CGT ATG CGC GT

CAC GCG CGT ATG CGC GT

CAC GCG CGT ATG CGC G

CAC GCG CGT ATG CGC

10 CAC GCG CGT ATG CGC

ACA CGC GCG TAT GCG CGT GTG G CA CGC GCG TAT GCG CGT GTG G A CGC GCG TAT GCG CGT GTG G CGC GCG TAT GCG CGT GTG G 15 GC GCG TAT GCG CGT GTG G C GCG TAT GCG CGT GTG G GCG TAT GCG CGT GTG G CG TAT GCG CGT GTG G G TAT GCG CGT GTG G 20 ACA CGC GCG TAT GCG CGT GTG ACA CGC GCG TAT GCG CGT GT ACA CGC GCG TAT GCG CGT G ACA CGC GCG TAT GCG CGT ACA CGC GCG TAT GCG CG 25 ACA CGC GCG TAT GCG C ACA CGC GCG TAT GCG ACA CGC GCG TAT GC

AC ACG CGC GTA TGC GCG TGT G
AC ACG CGC GTA TGC GCG TGT G
C ACG CGC GTA TGC GCG TGT G
ACG CGC GTA TGC GCG TGT G
CG CGC GTA TGC GCG TGT G
G CGC GTA TGC GCG TGT G
CGC GTA TGC GCG TGT G
GC GTA TGC GCG TGT G
CGC GTA TGC GCG TGT G
CGC GTA TGC GCG TGT G

CAC ACG CGC GTA TGC GCG TGT
CAC ACG CGC GTA TGC GCG TG
CAC ACG CGC GTA TGC GCG T
CAC ACG CGC GTA TGC GCG
CAC ACG CGC GTA TGC GC
CAC ACG CGC GTA TGC GC
CAC ACG CGC GTA TGC G
CAC ACG CGC GTA TGC
CAC ACG CGC GTA TGC

10 TCA CAC GCG CGT ATG CGC GTG T CA CAC GCG CGT ATG CGC GTG T A CAC GCG CGT ATG CGC GTG T CAC GCG CGT ATG CGC GTG T AC GCG CGT ATG CGC GTG T C GCG CGT ATG CGC GTG T 15 GCG CGT ATG CGC GTG T CG CGT ATG CGC GTG T G CGT ATG CGC GTG T TCA CAC GCG CGT ATG CGC GTG TCA CAC GCG CGT ATG CGC GT 20 TCA CAC GCG CGT ATG CGC G TCA CAC GCG CGT ATG CGC TCA CAC GCG CGT ATG CG TCA CAC GCG CGT ATG C TCA CAC GCG CGT ATG 25 TCA CAC GCG CGT AT

TTC ACA CGC GCG TAT GCG CGT G
TC ACA CGC GCG TAT GCG CGT G

30 C ACA CGC GCG TAT GCG CGT G
ACA CGC GCG TAT GCG CGT G
CA CGC GCG TAT GCG CGT G
A CGC GCG TAT GCG CGT G
CGC GCG TAT GCG CGT G
CGC GCG TAT GCG CGT G
C GCG TAT GCG CGT G
TTC ACA CGC GCG TAT GCG CGT
TTC ACA CGC GCG TAT GCG CG

TTC ACA CGC GCG TAT GCG C
TTC ACA CGC GCG TAT GCG
TTC ACA CGC GCG TAT GC
TTC ACA CGC GCG TAT G

TTC ACA CGC GCG TAT TC
TTC ACA CGC GCG TAT

ATT CAC ACG CGC GTA TGC GCG T TT CAC ACG CGC GTA TGC GCG T T CAC ACG CGC GTA TGC GCG T 10 CAC ACG CGC GTA TGC GCG T AC ACG CGC GTA TGC GCG T C ACG CGC GTA TGC GCG T ACG CGC GTA TGC GCG T CG CGC GTA TGC GCG T 15 G CGC GTA TGC GCG T ATT CAC ACG CGC GTA TGC GCG ATT CAC ACG CGC GTA TGC GC ATT CAC ACG CGC GTA TGC G ATT CAC ACG CGC GTA TGC 20 ATT CAC ACG CGC GTA TG ATT CAC ACG CGC GTA T ATT CAC ACG CGC GTA ATT CAC ACG CGC GT

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TAT TCA CAC GCG CGT ATG CGC G
AT TCA CAC GCG CGT ATG CGC G
T TCA CAC GCG CGT ATG CGC G
TCA CAC GCG CGT ATG CGC G

CA CAC GCG CGT ATG CGC G
A CAC GCG CGT ATG CGC G
CAC GCG CGT ATG CGC G
AC GCG CGT ATG CGC G
AC GCG CGT ATG CGC G
TAT TCA CAC GCG CGT ATG CGC
TAT TCA CAC GCG CGT ATG CG
TAT TCA CAC GCG CGT ATG C
TAT TCA CAC GCG CGT ATG C
TAT TCA CAC GCG CGT ATG C

TAT TCA CAC GCG CGT AT
TAT TCA CAC GCG CGT A
TAT TCA CAC GCG CGT
TAT TCA CAC GCG CG

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CTA TTC ACA CGC GCG TAT GCG C TA TTC ACA CGC GCG TAT GCG C A TTC ACA CGC GCG TAT GCG C TTC ACA CGC GCG TAT GCG C 10 TC ACA CGC GCG TAT GCG C C ACA CGC GCG TAT GCG C ACA CGC GCG TAT GCG C CA CGC GCG TAT GCG C A CGC GCG TAT GCG C CTA TTC ACA CGC GCG TAT GCG 15 CTA TTC ACA CGC GCG TAT GC CTA TTC ACA CGC GCG TAT G CTA TTC ACA CGC GCG TAT CTA TTC ACA CGC GCG TA CTA TTC ACA CGC GCG T 20 CTA TTC ACA CGC GCG CTA TTC ACA CGC GC

ACT ATT CAC ACG CGC GTA TGC G CT ATT CAC ACG CGC GTA TGC G 25 T ATT CAC ACG CGC GTA TGC G ATT CAC ACG CGC GTA TGC G TT CAC ACG CGC GTA TGC G T CAC ACG CGC GTA TGC G CAC ACG CGC GTA TGC G 30 AC ACG CGC GTA TGC G C ACG CGC GTA TGC G ACT ATT CAC ACG CGC GTA TGC ACT ATT CAC ACG CGC GTA TG ACT ATT CAC ACG CGC GTA T 35 ACT ATT CAC ACG CGC GTA ACT ATT CAC ACG CGC GT ACT ATT CAC ACG CGC G

ACT ATT CAC ACG CGC
ACT ATT CAC ACG CG

Examples of nucleobase sequences of peptide nucleic acid probes wherein the Qs of adjacent moieties are selected so as to be complementary to nucleobases within Position 1306 to 1322 shown in Figure 3 are the following ACA CCC ACC ACA AGG TGG ATG T
CA CCC ACC ACA AGG TGG ATG T
A CCC ACC ACA AGG TGG ATG T
CCC ACC ACA AGG TGG ATG T
CC ACC ACA AGG TGG ATG T
CACC ACA AGG TGG ATG T
ACC ACA AGG TGG ATG T
ACC ACA AGG TGG ATG T

CC **AC**A AG**G** TG**G** ATG T

15 C **AC**A AG**G** TG**G** ATG T

ACA CCC ACC ACA AGG TGG ATG
ACA CCC ACC ACA AGG TGG AT
ACA CCC ACC ACA AGG TGG A
ACA CCC ACC ACA AGG TGG

20 ACA CCC ACC ACA AGG TG
ACA CCC ACC ACA AGG T
ACA CCC ACC ACA AGG
ACA CCC ACC ACA AGG

CAC ACC CAC CAC AAG GTG GAT G 25 AC ACC CAC CAC AAG GTG GAT G C ACC CAC CAC AAG GTG GAT G ACC CAC CAC AAG GTG GAT G CC CAC CAC AAG GTG GAT G C CAC CAC AAG GTG GAT G 30 CAC CAC AAG GTG GAT G AC CAC AAG GTG GAT G C CAC AAG GTG GAT G CAC ACC CAC CAC AAG GTG GAT CAC ACC CAC CAC AAG GTG GA 35 CAC ACC CAC CAC AAG GTG G CAC ACC CAC CAC AAG GTG CAC ACC CAC CAC AAG GT

CAC ACC CAC CAC AAG G
CAC ACC CAC CAC AAG
CAC ACC CAC CAC AA

CCA CAC CCA CCA CAA GGT GGA T CA CAC CCA CCA CAA GGT GGA T A CAC CCA CCA CAA GGT GGA T CAC CCA CCA CAA GGT GGA T AC CCA CCA CAA GGT GGA T C CCA CCA CAA GGT GGA T 10 CCA CCA CAA GGT GGA T CA CCA CAA GGT GGA T A CC**A C**AA G**G**T G**G**A T CCA CAC CCA CCA CAA GGT GGA CCA CAC CCA CCA CAA GGT GG 15 CCA CAC CCA CCA CAA GGT G CCA CAC CCA CCA CAA GGT CCA CAC CCA CCA CAA GG CCA CAC CCA CCA CAA G CCA CAC CCA CCA CAA 20 CCA CAC CCA CCA CA

CCC ACA CCC ACC ACA AGG TGG A CC ACA CCC ACC ACA AGG TGG A C ACA CCC ACC ACA AGG TGG A 25 ACA CCC ACC ACA AGG TGG A CA CCC ACC ACA AGG TGG A A CCC ACC ACA AGG TGG A CCC ACC ACA AGG TGG A CC ACC **AC**A AG**G** TG**G** A 30 C ACC ACA AGG TGG A CCC ACA CCC ACC ACA AGG TGG CCC ACA CCC ACC ACA AGG TG CCC ACA CCC ACC ACA AGG T CCC ACA CCC ACC ACA AGG CCC ACA CCC ACC ACA AG CCC ACA CCC ACC ACA A CCC ACA CCC ACC ACA

• .

CCC ACA CCC ACC AC

ACC CAC ACC CAC CAC AAG GTG G CC CAC ACC CAC CAC AAG GTG G C CAC ACC CAC CAC AAG GTG G CAC ACC CAC CAC AAG GTG G AC ACC CAC CAC AAG GTG G C ACC CAC CAC AAG GTG G ACC CAC CAC AAG GTG G CC CAC CAC AAG GTG G 10 C CAC CAC AAG GTG G ACC CAC ACC CAC CAC AAG GTG ACC CAC ACC CAC CAC AAG GT ACC CAC ACC CAC CAC AAG G ACC CAC ACC CAC CAC AAG 15 ACC CAC ACC CAC CAC AA ACC CAC ACC CAC CAC A

> ACC CAC ACC CAC CAC ACC CAC ACC CAC CA

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Examples of nucleobase sequences of peptide nucleic acid probes wherein the Qs of adjacent moieties are selected so as to be complementary to nucleobases within Position 1621 to 1631 shown in Figure 3 are the following

GCC CCA GAA CTC CAC ACC CCC G

25 CC CCA GAA CTC CAC ACC CCC G

C CCA GAA CTC CAC ACC CCC G

CCA GAA CTC CAC ACC CCC G

CA GAA CTC CAC ACC CCC G

A GAA CTC CAC ACC CCC G

30 GAA CTC CAC ACC CCC G

AA CTC CAC ACC CCC G
A CTC CAC ACC CCC G

GCC CCA GAA CTC CAC ACC CCC

GCC CCA GAA CTC CAC ACC CC

35 GCC CCA GAA CTC CAC ACC C

GCC CCA GAA CTC CAC ACC

GCC CCA GAA CTC CAC AC

GCC CCA GAA CTC CAC A

GCC CCA GAA CTC CAC GCC CCA GAA CTC CA

CC CCA GAA CTC CAC ACC CCC
C CCA GAA CTC CAC ACC CC
CCA GAA CTC CAC ACC C
CCA GAA CTC CAC ACC
CA GAA CTC CAC ACC

10 ACT CCA CAC CCC CGA

Examples of nucleobase sequences of peptide nucleic acid probes wherein the Qs of adjacent moieties are selected so as to be complementary to nucleobases within Position 2401 to 2418 shown in Figure 4 are the following

15 CCG CCC CAA CTG GCG TCG AGG T
CG CCC CAA CTG GCG TCG AGG T
G CCC CAA CTG GCG TCG AGG T
CCC CAA CTG GCG TCG AGG T
CC CAA CTG GCG TCG AGG T

20 C CAA CTG GCG TCG AGG T
CAA CTG GCG TCG AGG T
AA CTG GCG TCG AGG T
A CTG GCG TCG AGG T
CCG CCC CAA CTG GCG TCG AGG

25 CCG CCC CAA CTG GCG TCG AG
CCG CCC CAA CTG GCG TCG A
CCG CCC CAA CTG GCG TCG
CCG CCC CAA CTG GCG TC
CCG CCC CAA CTG GCG T

30 CCG CCC CAA CTG GCG CCG CCC CAA CTG GC

TCC GCC CCA ACT GGC GTC GAG G
CC GCC CCA ACT GGC GTC GAG G

35 C GCC CCA ACT GGC GTC GAG G
GCC CCA ACT GGC GTC GAG G
CC CCA ACT GGC GTC GAG G
CC CCA ACT GGC GTC GAG G
C CCA ACT GGC GTC GAG G

CCA ACT GGC GTC GAG G
CA ACT GGC GTC GAG G
A ACT GGC GTC GAG G
TCC GCC CCA ACT GGC GTC GAG
TCC GCC CCA ACT GGC GTC GA
TCC GCC CCA ACT GGC GTC G
TCC GCC CCA ACT GGC GTC
TCC GCC CCA ACT GGC GTC
TCC GCC CCA ACT GGC GT
TCC GCC CCA ACT GGC GT
TCC GCC CCA ACT GGC G
TCC GCC CCA ACT GGC
TCC GCC CCA ACT GGC

CTC CGC CCC AAC TGG CGT CGA G TC CGC CCC AAC TGG CGT CGA G C CGC CCC AAC TGG CGT CGA G 15 CGC CCC AAC TGG CGT CGA G GC CCC AAC TGG CGT CGA G C CCC AAC TGG CGT CGA G CCC AAC TGG CGT CGA G CC AAC TGG CGT CGA G 20 C AAC TGG CGT CGA G CTC CGC CCC AAC TGG CGT CGA CTC CGC CCC AAC TGG CGT CG CTC CGC CCC AAC TGG CGT C CTC CGC CCC AAC TGG CGT 25 CTC CGC CCC AAC TGG CG CTC CGC CCC AAC TGG C CTC CGC CCC AAC TGG CTC CGC CCC AAC TG

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GCC CCA ACT GGC GTC

ACT CCG CCC CAA CTG GCG TCG A
CT CCG CCC CAA CTG GCG TCG A

T CCG CCC CAA CTG GCG TCG A
CCG CCC CAA CTG GCG TCG A
CG CCC CAA CTG GCG TCG A
G CCC CAA CTG GCG TCG A

CCC CAA CTG GCG TCG A
CC CAA CTG GCG TCG A
C CAA CTG GCG TCG A
ACT CCG CCC CAA CTG GCG TCG
ACT CCG CCC CAA CTG GCG TC
ACT CCG CCC CAA CTG GCG T
ACT CCG CCC CAA CTG GCG
ACT CCG CCC CAA CTG GCG
ACT CCG CCC CAA CTG GC
ACT CCG CCC CAA CTG GC
ACT CCG CCC CAA CTG GC
ACT CCG CCC CAA CTG
ACT CCG CCC CAA CTG

Examples of nucleobase sequences of peptide nucleic acid probes wherein the Qs of adjacent moieties are selected so as to be complementary to nucleobases within Position 2455 to 2486

shown in Figure 4 are the following

CTA AAC CCG ATT CAG GGT TCG A
CTA AAC CCG ATT CAG GGT TCG
CTA AAC CCG ATT CAG GGT TC

CTA AAC CCG ATT CAG GGT T CTA AAC CCG ATT CAG GGT

CTA AAC CCG ATT CAG GG

CTA AAC CCG ATT CAG G

CTA AAC CCG ATT CAG
CTA AAC CCG ATT CA

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CAG GGT TCG AGG TTA GAT GCC C
AG GGT TCG AGG TTA GAT GCC C
G GGT TCG AGG TTA GAT GCC C
GGT TCG AGG TTA GAT GCC C
GGT TCG AGG TTA GAT GCC C
T TCG AGG TTA GAT GCC C
TCG AGG TTA GAT GCC C
CG AGG TTA GAT GCC C
G AGG TTA GAT GCC C
CAG GGT TCG AGG TTA GAT GCC
CAG GGT TCG AGG TTA GAT GCC
CAG GGT TCG AGG TTA GAT GC
CAG GGT TCG AGG TTA GAT GC

CAG GGT TCG AGG TTA GAT

CAG GGT TCG AGG TTA GA

TCA GGG TTC GAG GTT AGA TGC C CA GGG TTC GAG GTT AGA TGC C A GGG TTC GAG GTT AGA TGC C GGG TTC GAG GTT AGA TGC C GG TTC GAG GTT AGA TGC C G TTC GAG GTT AGA TGC C TTC GAG GTT AGA TGC C TC GAG GTT AGA TGC C 10 C GAG GTT AGA TGC C TTC GAG GTT AGA TGC TCA GGG TTC GAG GTT AGA TGC TCA GGG TTC GAG GTT AGA TG TCA GGG TTC GAG GTT AGA T 15 TCA GGG TTC GAG GTT AGA

TTC AGG GTT CGA GGT TAG ATG C

TC AGG GTT CGA GGT TAG ATG C

20 C AGG GTT CGA GGT TAG ATG C

AGG GTT CGA GGT TAG ATG C

GG GTT CGA GGT TAG ATG C

G GTT CGA GGT TAG ATG C

GTT CGA GGT TAG ATG C

TT CGA GGT TAG ATG C

TT CGA GGT TAG ATG C

TTC AGG GTT CGA GGT TAG ATG

TTC AGG GTT CGA GGT TAG ATG

TTC AGG GTT CGA GGT TAG AT

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CTG TCC CTA AAC CCG
GTC CCT AAA CCC GAT

Examples of nucleobase sequences of peptide nucleic acid probes wherein the Qs of adjacent moieties are selected so as to be complementary to nucleobases within Position 3094 to 3103 shown in Figure 4 are the following CAG GTC TGA CCT ATT GAA CCC G
AG GTC TGA CCT ATT GAA CCC G

G GTC TGA CCT ATT GAA CCC G GTC TGA CCT ATT GAA CCC G TC TGA CCT ATT GAA CCC G C TGA CCT ATT GAA CCC G TGA CCT ATT GAA CCC G GA CCT ATT GAA CCC G A CCT ATT GAA CCC G CAG GTC TGA CCT ATT GAA CCC CAG GTC TGA CCT ATT GAA CC CAG GTC TGA CCT ATT GAA C 10 CAG GTC TGA CCT ATT GAA CAG GTC TGA CCT ATT GA CAG GTC TGA CCT ATT G CAG GTC TGA CCT ATT CAG GTC TGA CCT AT 15

> AG GTC TGA CCT ATT GAA CCC G GTC TGA CCT ATT GAA CC GTC TGA CCT ATT GAA C TC TGA CCT ATT GAA

EXAMPLES

EXAMPLE 1

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In situ hybridisation to fixed bacterial cells

To test the ability of the peptide nucleic acid probes to detect MTC and not MAC or Neisseria gonorrhoeae, fluorescence *in situ* hybridisation (FISH) was performed on fixed bacterial cells using fluorescein labelled probes as shown below. It was shown that these probes did not hybridise to M. avium, M. intracellulare, or N. gonorrhoeae.

Preparation of bacterial slides

M. bovis BCG (Statens Seruminstitut, Denmark, Catalogue number 2645), M. avium (Statens Seruminstitut, Denmark, Laboratory number 3716 (E37978)), and M. intracellulare (Statens Seruminstitut, Laboratory number 3717 (E39562)) were grown in Dubos medium (Statens Seruminstitut, Denmark) or on Löwenstein-Jensen medium (Statens Seruminstitut, Denmark) at 37 °C. N. gonorrhoeae was grown on chocolate agar at 37 °C with additional 5% CO₂.

Bacterial smears were prepared on test slides according to standard procedures. The smears were air-dried followed by flame fixation.

FISH on bacterial slides

- 5 The following procedure was performed.
 - 1. The slide is immersed in 80% ethanol for 15 minutes, subsequently rinsed with water and air-dried.
 - 2. The bacterial slide is covered with a hybridisation solution containing the probe in question at a concentration of 250 nM.
- 10 3. The slide is incubated in a humid incubation chamber at 45 °C for 90 minutes.
 - 4. The slide is washed 25 minutes in TBS-buffer, pH 10 at 45 °C, followed by 30 seconds in water.
 - 5. The slide is dried and mounted (DAKO Fluorescence Mounting Medium or equivalent).
- 15 The following hybridisation solutions was used:

Hybridisation

10 mM NaCl

solution

10% Dextran sulphate

30% formamide

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0.1% Triton X-100®

50 mM Tris-HCl, pH 7.6

50 mM EDTA

0.1% sodium pyrophosphate

0.2% polyvinylpyrrolidone

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0.2% Ficol

TBS buffer

10 mM sodium phosphate, pH 10

145 mM NaCl

30 All solutions are made RNase free following standard procedures.

The following peptide nucleic acid probe was used

Lys(Flu)-Lys(Flu)-TTC GAG GTT AGA TGC-NH₂ Lys(Flu)-Lys(Flu)-ACT CCA CAC CCC CGA-NH₂ **OK 306**

OK 309

wherein Flu denotes a fluorescein isothiocyanate label or a 5-(and 6)-carboxyfluorescein label, and Lys(Flu)-Lys(Flu) denotes a peptide label ("zipper") with two Flu labels attached. The

results are shown in Table 1.

TABLE 1

Probe OK 306	FISH
M. bovis BCG	positive
M. avium	negative
M. intracellulare	negative
N. gonorrhoeae	negative

Probe OK 309	FISH
M. bovis BCG	positive
M. avium	negative
M. intracellulare	negative
N. gonorrhoeae	negative

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EXAMPLE 2

Test in dot blots

To further test the ability of the peptide nucleic acid probes to detect MTC and not MAC or E. coli, dot blot tests were carried out.

M. bovis BCG (Statens Seruminstitut Catalogue number 2645) and M. intracellulare (Statens Seruminstitut, Denmark Laboratory number 3713 (E39562)) were grown in Dubos medium (Statens Seruminstitut, Denmark) or on Löwenstein-Jensen medium (Statens Seruminstitut, Denmark) at 37 °C.

RNA was isolated from the bacterial cells by use of TRI-reagent (Sigma) following manufacture's directions. E. coli rRNA was purchased from Boehringer Mannheim, Germany.

20 The following nucleic acid probes were used.

Lys(Flu)-Lys(Flu)-CTG TCC CTA AAC CCG-NH₂	OK 305
Lys(Flu)-Lys(Flu)-GTC CCT AAA CCC GAT-NH₂	OK 307
Lys(Flu)-Lys(Flu)-ACT CCA CAC CCC CGA-NH₂	OK 309

wherein Flu denotes a fluorescein isothiocyanate label or a 5-(and 6)-carboxyfluorescein label, and Lys(Flu)-Lys(Flu) denotes a peptide label ("zipper") consisting of 2 amino acids, respectively, with two Flu labels attached.

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Preparation of dot blots

The following buffers were used:

20 x SSPE buffer 3 M NaCl

10 0.2 M PO₄³⁻

0.02 M EDTA

pH 7.4

TST buffer 0.05 M Tris/HCI

0.5 M NaCl

0.5% Tween 20®

pH 9.0

200 ng M. bovis RNA, M. intracellulare RNA and E. coli rRNA were dotted onto membranes (Schleich & Schuel, NY 13 N), and the membranes were dried and fixed under UV light for 2 minutes. Each of the probes (70 nM probe in hybridisation solution (hybridisation solution without Triton X-100® and with the exception that formamide was substituted with 50% glycerol)) were added to the membrane. Hybridisation was continued for 1.5 hours at 55 °C. The membranes were rinsed 2 times for 15 minutes in 2 × SSPE buffer containing 0.1% SDS at ambient temperature, and subsequently 2 times for 15 minutes in 0.1 × SSPE buffer containing 0.1% SDS at 55 °C or at 65 °C (see Table 2). The membrane was blocked with 0.5% casein dissolved in 0.05M Tris/HCl and 0.5 M NaCl with pH 9.0. Thereafter, the membranes were incubated for 1 hour with rabbit-anti FITC antibody labelled with AP (DAKO K0046 vial A) diluted 1:2000 in 0.5% casein dissolved in 0.05M Tris/HCl and 0.5 M NaCl with pH 9.0. After incubation, the membranes were washed 3 times 5 minutes with TST at ambient temperature. Bound probes were visualised following standard procedures using BCIP/NBT, and the visualisation was stopped by incubation for 10 minutes with 10 mM EDTA. The blot was dried at 50 °C.

The results are given in Table 2 below.

TABLE 2

	E. coli rRN	ΝA	M. bovis BCG RNA		M. intracellulare RNA	
Probe	55 °C	65 °C	55 °C	65 °C	55 °C	65 °C
OK 305	negative	negative	positive	positive	negative	weak
OK 307	negative	negative	positive	positive	negative	weak
OK 309	negative	negative	positive	positive	negative	weak

CLAIMS

1. A peptide nucleic acid probe for detecting mycobacteria of the Mycobacterium tuberculosis Complex (MTC) in a sample, which probe comprises from 10 to 30 polymerised moieties of formula (I)

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wherein each X and Y independently designate O or S, each Z independently designates O, S, NR^1 , or $C(R^1)_2$, wherein each R^1 independently designate H, C_{1-6} alkyl, C_{1-6} alkenyl, C_{1-6} alkynyl, each R^2 , R^3 and R^4 designate independently H, the side chain of a naturally occurring amino acid, the side chain of a non-naturally occurring nucleobase, C_{1-4} alkyl, C_{1-4} alkenyl or C_{1-4} alkynyl, or a functional group, each Q independently designates a naturally occurring nucleobase, a non-naturally occurring nucleobase, an intercalator, a nucleobase-binding group, a label or H,

with the proviso that the Qs of adjacent moieties are selected so as to form a sequence of which a subsequence includes at least one nucleobase complementary to a nucleobase of M. tuberculosis 23S rRNA that differs from the corresponding nucleobase of M. avium located within the following domains

Position 326 to Position 359 in Figure 1, or Position 635 to Position 658 in Figure 1, or Position 761 to Position 787 in Figure 2, or Position 1306 to Position 1322 in Figure 3, or Position 1621 to Position 1631 in Figure 3, or Position 2401 to Position 2418 in Figure 4, or Position 2455 to Position 2486 in Figure 4, or Position 3094 to Position 3103 in Figure 4,

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and further with the proviso that the probe comprising such subsequence is able to form hybrids with target sequences in 23S rRNA of said mycobacteria.

2. A peptide nucleic acid probe according to claim 1 of formula (II), (III), or (IV)

$$\sum_{\mathbb{R}^3} \mathbb{R}^3$$
 (IIII)

$$\sum_{N} N$$
 (IV)

wherein Z, R2, R3, and R4, and Q is as defined in claim 1.

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- 3. A peptide nucleic acid probe according to claim 1 or 2, wherein Z is NH, NCH₃ or O, each R², R³ and R⁴ independently designate the side chain of a naturally occurring nucleobase, the side chain of a non-naturally occurring nucleobase, or C₁₋₄ alkyl, and each Q is a naturally occurring nucleobase or a non-naturally occurring nucleobase with the provisos defined in claim 1.
 - 4. A peptide nucleic acid probe according to anyone of claims 1 to 3, wherein Z is NH or O, and R² is H or the side chain of Ala, Asp, Cys, Glu, His, HomoCys, Lys, Orn, Ser or Thr, and Q is a nucleobase selected from thymine, adenine, cytosine, guanine, uracil, iso-C, iso-G and 2,6-diaminopurine with the provisos defined in claim 1.
 - 5. A peptide nucleic acid probe according to anyone of claims 1 to 4 of formula (V)

wherein R^4 is H or the side chain of Ala, Asp, Cys, Glu, His, HomoCys, Lys, Orn, Ser or Thr, and Q is as defined in claim 4 with the provisos defined in claim 1.

- 6. A peptide nucleic acid probe according to anyone of claims 1 to 5 further comprising one or more labels which may be mutually identical or different, and/or one or more linkers which may be mutually identical or different with the provisos defined in claim 1.
- 7. Method for detecting mycobacteria of the Mycobacterium tuberculosis Complex in a sample comprising
- (1) contacting any rRNA optionally present in said sample with one or more peptide nucleic acid probes according to anyone of claims 1 to 6 under conditions, whereby hybrids between said probe(s) and said rRNA are formed, and
 - (2) observing or measuring said hybridisation, and relating said observation or measurement to the presence of mycobacteria of the Mycobacterium tuberculosis Complex in said sample.
 - 8. Method according to claim 7, c h a r a c t e r i s e d in that the hybrids are captured on a solid phase before measuring the
 - 9. Method according to claim 7, c h a r a c t e r i s e d in that a peptide nucleic acid probe according to anyone of claims 1 to 6 are used for capturing the hybrids.
- 10. A method according to anyone of claims 7 to 9, c h a r a c t e r i s e d in that a signal amplifying system is used for measuring the resulting hybridisation.
 - 11. Kit for detecting mycobacteria of the Mycobacterium tuberculosis Complex, c h a r a c t e r i s e d in that said kit comprises at least one peptide nucleic acid probe according to anyone of claims 1 to 6, and a detection system with at least one detecting reagent.
- 12. Kit according to claim 11,35 characterised in that it further comprises a solid phase capture system.

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extent of hybridisation.

ABSTRACT

NOVEL PROBES FOR THE DETECTION OF MYCOBACTERIA OF THE MYCOBACTERIUM TUBERCULOSIS COMPLEX

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Novel hybridisation assay probes for detecting mycobacteria of the Mycobacterium tuberculosis Complex (MTC) are provided. The probes detect 23S rRNA of MTC. Such probes are capable of detecting the organisms in test samples, e.g. expectorates, sputum, aspirates, urine, blood and tissue sections, food, soil and water.

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		1/-	∓	320	
	290	300	310		
1251	ACGCATGGGTAACC	GGGTAGGGGTT	TGTGTGCG	GGGTTG 1	M.tuberculosis
579	A FIND CONTROL TO A A C C	CCCTAGGGGTT	STGTGTGCG	المالك	M.avium
665		\Box	TCTCTGCG	. فالمالة	M. DIITET
591			ヹ゙ヸ゙゙゙゙゚゚ゟヸ゙゙゙゚ヹヸ゙゙゚ヸ゙゚゚゚ヹヸヸヸヸヸヸヸヸヸヸヸヸヸヸ	71(G1"1"G	M. lebrae
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366	A COCA MOCOCOMO CO	いことには、これには、これには、これには、これには、これには、これには、これには、これ	7116116116	GGGIIG	M. gaber -
309	ACGCATGGGTAACC	GGGTAGGGGTT	GTGTGTGCG	GGGTTG	M.kansasi
303		340	350	360	•
	330				s tubemoulogie
1291	TGGGAG-GATATGT	CTCAGCGCTAC	CCGCTGAG	A-GGCA	M.tuberculosis
619	magaz Frida y Ta Ta	r CTCACC $\overline{\Pi}$ CTAC	CTGGCTGAG	G-GGIA	M.avium
705		$^{\circ}$ CAUCAMCGTCCG	CCGCGAIG	CCACIA	M. PHICE
631			CYTOCACA (11/11/12/DC)	[M. IEDIAC
619	TO CONTROL TO TO	CTCAGCTCTAC	CTGGCTGAG	G - GGIA	M. paracuberc.
406		"C" " " " " " " " " " " " " " " " " " "	CCGCTGAG	G-66CA	M.gaberr
349	TGGGATCGATACG	CTCAGCTCTAC	CCGGCTGAG	G-GGCA	M.kansasi
2.2		380	390	400	
	370			T T T T T T T T T T T T T T T T T T T	M tuberculosis
1329	GTCAGAAAGTGTC	TGGTTAGCGGA	AGTGGCCTG	GGATGG	M.tuberculosis
658	as a samemer	マͲイイイͲͲϦ ႺイイႺႺϪ	Δ CTCCCTC	JUMPH	M.avium
744		っかくしかがく へんかごり	\mathbf{z}	UULA:	M. Durer
670		<u>ነውረረውውን ሮሮሮርን</u>	$\Delta \Delta T = C + C + C + C + C + C + C + C + C + C$	シシょみむむ	M. Tebrae
658		?ͲϤϹͲͲϪϹϹϢϢ	$A(\frac{1}{2}) \cup \{0\} \cup \{1\}$	DUDAUU.	M. paracaser.
445		~中へ~中中カル~~(ころ)	Δ GTGGCCTU	JULA JUG	M. gastri
388	GTCAGAAAGTGTC	GTGGTTAACGGA	AGTGGCCT	GGATGG	M.Kansası
		* * *	* *		
	610	620	630	640)
				THE COMME	M tuberculosis
1568	ACCTGAAACCGTG'	rgcctacaatcc	GTCAGAGCC	TCCTTT	M.tuberculosis
896	A COMON NA COCOTO	rCCCTACAATCC	GTCAGAGCC	TUCTICG	M.avium
978	ACCTGAAACCGTG'	rgcctacaatcc	GTCAAAGCC	CICICI	M. Ponter
909	ACCTGAAACCGTG'	TGCCTACAATCC	GTCAGAGCC	TGITIG	M. reprae
896	አ ረረጥሮ እ እ እ ሮሮርጥር'	TGCCTACAATCC	GTCAGAGC(Trecrice	M.paracuberc.
684	A COMON N N COOPE	$_{TCCCTTDCDDTCC}$	'GTCAGAGC	MITTIN	M. yastıı
627	ACCTGAAACCGTG	TGCCTACAATCC	CGTCAGAGC	CTTTCG	M.Kansası
	650	660	670	68	
			TO COMPO COM	POTCAAC	M.tuberculosis
1608	TCCTCTCCGGAGG	AGGGTGGTGAT	GCGTGCCT GCGTGCCT	\mathbf{r}	M avium
936	T	GGGGTGATC	GCGTGCCT	\mathbf{n}	M phlei
1018	3 TGT	AGTGGGGTGAT	-GCGTGCCT	₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽	M lenrae
949	T	GGGGTGAT		╨╨╨╓ <i>╸</i> ┸┸┸╚╇╋╚	M.leprae
936	T	GGGGTGAT	~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	1110WW	M.paratuberc.
724	T	GGGGTGAT		TTTGAAG	M.gastri
667	T		_	TITGAAG	M.kansasi
		Figu	re 1		

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-	730	740	750	760	
1001 1088 1014 1001	GTGTGGGGTAGCCGC GTGTGGGGGTAGCCGC GTGTGGGGGTAGCCGC GTGTGGGGGTAGCCGC GTGTGGGGGTAGCCGC GAGCGGGGGTAGCCGC	ZAGCGAAAGCC ZAGCGAAAGCC ZAGCGAAAGCC ZAGCGAAAGCC	SAGTCTGAATAC SAGTCTGAATAC SAGTCTGAATAC SAGTCTGAATAC	GGGCG M.aviu GGGCG M.phle: GGGCG M.lepra GGGCG M.para GGGCG M.gast:	" ae tuberc. ri
1728 1041	770 ACCCACACGCGCATA CATCQCCTTTC TATCCAACCTGTTC TATCACGTGT CATCACGTGT TATCACGCGT TATCACGCGT	780 ACGCGCGTGTG AGGTTGGTG AGCGTGTG AGCGTGTG AGCGTGTG AGCGTGTG AGCGTGTG AGCGTGTG	790 GAATAGTGGCGT TAGTGGCGT TAGTGGCGT TAGTGGCGT TAGTGGCGT TAGTGGCGT TAGTGGCGT TAGTGGCGT	800 TGTTC M.tube: TGTTC M.avius TGTTC M.phle TGTTC M.lepr TGTTC M.para TGTTC M.gast TGTTC M.kans	rculosis m i ae tuberc. ri asi
1768 1070 1164 1086 1070 861 804	TGGACCCGAAGCGG TGGACCCGAAGCGG TGGACCCGAAGCGG	AGTGATCTAC AGTGATCTAC AGTGATCTAC AGTGATCTAC	CCATGGCCAGG CCATGGCCAGG CCATGGCCAGG CCATGGCCAGG	GTGAA M.phle GTGAA M.lepi GTGAA M.para GTGAA M.gast	ei rae atuberc. tri

	200	1300		1320	
	1	TOO CA CATTO	ACC - TTGTG	GTG 1	M.tuberculosis
2246 GCACACCG	CCGAAGCCG	GGCACATEC	ACC-TTACG	GTG 1	M.avium
1549 GCACACCG 1643 GCACACCG	CCGAAGCCG	GGCACATHC	GCC-UTTGTG	GCT :	M.phlei
1643 GCACACCG	CCGAAGCCG	CGGCAGATCE		GTG	M.leprae
1643 GCACACCG 1565 GCACACCG	CCGAAGCCG	CGGCACATIL	ACCITCIAGO	GTG	M paratuberc.
1565 GCACACCG 1549 GCACACCG	CCGAAGCCG	CGGCACATIC	ALCITICATION	GT∏	M.gastri
1339 GCACACCG 1282 GCACACCG	CCGAAGCCG	CGACAACCG	A	01 <u> </u>	
1	.330	1340	1350	1360	
	ma cccca cc	GTCCCTCAT'	TCAGCGAAGCC	ACC	M.tuberculosis M.avium
2284 <u>GG</u> TGTGGG 1588 GATGTGGG	TAGGGGAGC	GTCCCTCAT'	TCAGCGAAGCT	-CC	M.avium
1681 GGTGTGGG	TAGGGGAGC	CTTCCTCAT'	TCAGCGAAGCC	TICC	M.leprae
1605 GATGTGGG	TAGGGGAGC	CTCCCCCAT	TCAGCGAAGCI	-cc	M.paratuberc. M.gastri
1588 GATGTGGG	TAGGGGAGC	CTCCCTCAT	TCAGCGAAGCC	GCC	M.gastri
1371 TGGG	TAGGGGAGC	CTCCCTCAT	TCAGCGAAGC	dcc	M.kansasi
1314 TGGG	JAGGGGAGC	.G.CC.CA.			
		***		164	0
:	1610	1620	1630		
AFGA AMONG TO	CCCTTCGGG	GG-TGTGGA	GTTCTGGGGCT	GCG	M.tuberculosis M.avium
1865 AGCAT-TO 1960 ATCAT 1884 AGCATATO		GGTGAC	GGTTGGGGGCT	rgcg	M.phlei
1960 ATCATE		GG TATGGA	GGTTGGGGGCT	rgcg	M.leprae
1865 AGCAT-TO	CCCTTCGG	GGA - GTGGA	GTCTGGGGC	rgcg	M.gastri
1646 ATCAC-TO 1589 ATCAC-TO		GGC-GTGGA	GTCTGGGGC	rgcg	M.kansasi
1589 ATCAC-TO			1670	168	0
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2601 TGGGAAC	TTCGCTGGT	AGTAGTCA	-AGCGAAGGG	-GTG	M.tuberculosis M.avium
1903 TGGGAGG	CG-GTGGGT	AGTAGTC	A-AGCGATGGG	-GTG	M.phlei
			a according		
TAZA IGGGWWC	$\operatorname{TTCGTTGGT}$				
		A GTAGTCA	A-AGCGATGGG A-AGCZATGGG		
1903 TGGGACC	TTCGCTGGT.	A GTAGTCA A GTAGTCA A - CTAGTCA	1-AGCGATIGGG 1-AGCGATIGGG 1-AGCGATIGGG	-GTG	M.gastri
1903 TGGGACC	TTCGCTGGT.	A GTAGTCA A GTAGTCA A - CTAGTCA		-GTG	M.gastri

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	2	2370	2380	2390	2400	
		LCTTTTCTC	TACCATACCT	GGGAGACTG	TGAAAC N	1.tuberculosis
3307	TCGGTACG		TAGGATAGGT	GGGAGACTÍ	TGAAGC N	1.avium
			アメクロスアメロロア	CCCACACIL	TGAAGC I	1. PIII-CI
2630	TCGGTGCG	GTTTGTG	TAGGATAGGI	CCCACACTT	TGAAGC 1	M.paratuberc.
	TCGGTACG	GTTTGTG	TAGGATAGGT	GGGAGACIE	ייים	M.gastri
1910		· · ·		CCCNCNCTC	- בתכמממר 1	M kansasi
2334	TCGGTACC	GTTTGTG	TAGGATAGGT		2440	
	:	2410	2420	2430	2440	
2247	CTCCA CCC	CACTTGG	GGGGAGTCG	TTGTTGAAA	TACCAC I	M.tuberculosis
3347			momeon cree	TTGTTGAAB	TACCAC I	1.avium
2647		~~~ ammida		TTTCTCAAA	AIACCAC A	71 - P11± C ±
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2670	IIITCGACGC	LIAGI IGG	TCTCGACTCC	TTGTTGAA	ATACCAC I	M.paratuberc.
2647						
1910		~~> ~~	GGTGGAGTCG	TTGTTGAA	ATACCAC	M.kansasi
2374	CTCAACG	CCAGTTGG	GG I GGAGI CC		2480	\
		2450	2460	2470		
2207	TOTONTO	<u>፲</u> ፻፮ ምጥርር	CATCTAACCI	CGAACCCT	GAATCGG	M.tuberculosis
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2414	. TCTGATC	GIATIGG			050	_
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2427	CTTTACC	CACAGTGO	CTGGCGGGT	GTTTAACT	GGGGCGG	M.tuberculosis
3427		へ 入 へ 入	$^{\circ}$ CTGGCGGGT $)$	AGTTTAACT	<i>555555</i>	M. avium
	COMPICIA CO	ペカペカでTCC	тстсстссст	AGTTTAACT		M. Durer
2819		~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~	~~~~~~~~~~~	יויי אם מיויידידים מ	しんけんしんしゅん	M. Tebrac
2750	GTTTAGG	GACAGIG(CTGGCGGGT	AGTTTAACT	GGGGCGG	M.paratuberc.
						rr. qubur-
1910) - cmmGa/GC	<u> </u>	CCTGGCGGGT.	AGTTTAACT	GGGGCGG	M.kansasi
2454	I GIII LICALIG	GACAGIG	C16666661			
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		3090	3100	3110	312	
				A CA CCTCCA	ACCTCAG	M.tuberculosis
402	6 GC-AGAA	CACGGG	TCAATAGGIC	AGACCI GGA	ACCTCAG	M avium
332	5 GC-AGA	CACGGGA	TTGATAGGC		MGC1CVG	M. phlei
341	8 GC-AGA	CACGGGA	TCGATAGACC	AGACCIGGE	AGGCACAG	M.leprae
						M. TEDIAC
332	5 GC-AGA	CACGGGA	TTGATAGGCC	AGACCTGGA	MGCICHA	M.paratuberc. M.gastri
305	3 GC-AGA	ACACGGGT	TCGATAGGGC	AGACCTGGA	AAGCTCAG	M.kansasi
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